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(54) Title: THREE-DIMENSIONAL STRUCTURE OF DIPEPTIDYL PEPTIDASE IV

(57) Abstract: A crystal of a dipeptidyl peptidase IV; a three-dimensional structural coordinate of the dipeptidyl peptidase IV; a method for obtaining a three-dimensional coordinate of a homolog protein of the dipeptidyl peptidase IV; a method for obtaining a three-dimensional structural coordinate of a crystal of a complex of the dipeptidyl peptidase IV and a effector of the dipeptidyl peptidase IV; a method for identifying pharmacophore of the effector of the dipeptidyl peptidase IV; a method for designing, identifying, evaluating or searching; the effector; and a program and a medium therefor for use of the three-dimensional structural coordinate.

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DESCRIPTION

THREE-DIMENSIONAL STRUCTURE OF DIPEPTIDYL PEPTIDASE IV

5 TECHNICAL FIELD

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The present invention relates to a crystal and a three-dimensional structural coordinate of a dipeptidyl peptidase IV, and an application thereof. More specifically, the present invention relates to a crystal and a threedimensional structural coordinate, a method for obtaining a three-dimensional structural coordinate of a homolog protein of a dipeptidyl peptidase IV, a method for obtaining a three-dimensional structural coordinate of a crystal of a complex of a dipeptidyl peptidase IV with an effector (e.g. inhibitor) of the dipeptidyl peptidase IV, a method for identifying a pharmacophore of an effector (e.g. inhibitor) of for the dipeptidyl peptidase IV, a method for identifying sites affecting the activity of the dipeptidyl peptidase IV, a method for designing, identifying, evaluating or searching an effector (e.g. inhibitor) of the dipeptidyl peptidase IV, and a program and a medium therefor for use of the three-dimensional structural coordinate, which are useful in the development of an effector (e.g. inhibitor) of the dipeptidyl peptidase IV, useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like; and an effector (e.g. inhibitor) of the dipeptidyl peptidase IV useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like.

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BACKGROUND ART

Dipeptidyl peptidase IV (hereinafter also referred to as DPPIV) is a cell membrane protein, which has been found in epithelial cell of small intestine, prostate gland, renal tubule, biliary tract and the like, activated T-cell, B-cell, NK-cell and the like. In the DPPIV, deduced active sites of DPPIV in the C-terminal side are located in extracellular portions and those in the N-terminal side are located in cytoplasm in a living body. Also, there has been suggested the relationship of the above-mentioned DPPIV with the activities of various cytokines such as interleukin-1β, interleukin-2, interleukin-3, interleukin-5, interleukin-6, interleukin-13, tumor necrosis factor-β and the like, and activities of various chemokines such as RANTES and the like in immune system [Rinsho Menneki (Clinical Immunology), 34, Revised and Enlarged Edition 19, 45-53, published by Kagaku Hyoronsha (2000), and the like].

As to the dipeptidyl peptidase IV, it has been shown that some amino acid residues can be involved in exhibition of the activity of the dipeptidyl peptidase IV by experiments such as biochemical experiments using inhibitors, experiments using mutants produced by site-directed mutagenesis [for example, see Misumi et al, Biochim. Biophys. Acta, 1131, 333-336 (1992), Ogata et al,

20 Biochemistry, 31, 2582-2587 (1992) and the like].

However, it is difficult to know the three-dimensional structures for active sites from the information. Therefore, it is presently difficult to obtain the three-dimensional structural information for identifying, searching, evaluating or designing an interaction of the dipeptidyl peptidase IV and a compound that acts with the dipeptidyl peptidase IV on the level of three-dimensional structure and a

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novel compound capable of binding with and acting on the dipeptidyl peptidase IV.

DISCLOSURE OF INVENTION

A first object of the present invention is to provide a crystal of a dipeptidyl peptidase IV, which is useful for providing a three-dimensional structural coordinate as the information for designing, identifying, evaluating or searching an effector (e.g. inhibitor) of the dipeptidyl peptidase IV useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like. A second object of the present invention is to provide a three-dimensional structural coordinate of the crystal. which can provide the information for designing, identifying, evaluating or searching an effector (e.g. inhibitor) of the dipeptidyl peptidase IV useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like. A third object of the present invention is to provide a method for obtaining a three-dimensional structural coordinate of a homolog protein of the dipeptidyl peptidase IV, whereby refinement of a three-dimensional structural coordinate of a homolog protein of the dipeptidyl peptidase IV can be more readily performed. Furthermore, a fourth object of the present invention is to provide a method for obtaining a three-dimensional structural coordinate of a crystal of a complex of a dipeptidyl peptidase IV and an effector (e.g. inhibitor) of the dipeptidyl peptidase IV, which can provide the information for designing, identifying, evaluating or searching an

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effector (e.g. inhibitor) of the dipeptidyl peptidase IV which is useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and is excellent in avidity, biological activity, biological stability, absorbency to a living body, and which can favorably act on the dipeptidyl peptidase IV. A fifth object of the present invention is to provide a method for identifying a pharmacophore of the dipeptidyl peptidase IV and the effector (e.g. inhibitor) of the dipeptidyl peptidase IV, which can provide the information for designing, identifying, evaluating or searching an effector (e.g. inhibitor) of the dipeptidyl peptidase IV useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and is excellent in avidity, biological activity, biological stability, absorbency in a living body, and which can be favorably act on the dipeptidyl peptidase IV. A sixth object of the present invention is to provide a method for designing, identifying, evaluating or searching the effector (e.g. inhibitor) of the dipeptidyl peptidase IV, which can logically and conveniently provide the effector (e.g. inhibitor) of the dipeptidyl peptidase IV useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and is excellent in avidity, biological activity, biological stability, absorbency in a living body (in vivo), and which can be favorably act on the dipeptidyl peptidase IV. A seventh object of the present invention is to provide the effector (e.g. inhibitor) of the dipeptidyl peptidase IV useful as a modulatory agent of immune

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response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like. An eighth object of the present invention is to provide a program and a medium therefor, which can rapidly and conveniently perform design, identification, evaluation or search of the effector (e.g. inhibitor) of the dipeptidyl peptidase IV.

Concretely, the present invention relates to:

- [1] a crystal of a dipeptidyl peptidase IV, having characteristics sufficient to ensure a resolution capable of analyzing its three-dimensional structure up to the side chain level by X-ray crystallographic structural analysis;
- [2] the crystal according to the above [1], wherein the dipeptidyl peptidase IV is a soluble polypeptide comprising a region located at extramembrane in a full-length human dipeptidyl peptidase IV;
- [3] the crystal according to the above [1] or [2], wherein the dipeptidyl peptidase IV is a polypeptide having an amino acid sequence in which a transmembrane region is deleted from the amino acid sequence of SEQ ID NO: 2, and a tag peptide is optionally added to a C-terminal side or N-terminal side thereof;
- [4] the crystal according to any one of the above [1] to [3], wherein the crystal has a space group of $P2_12_12_1$, and a lattice constant of the unit cell of $|a| = 118.0 \pm 5.0$ Å, $|b| = 125.9 \pm 5.0$ Å, $|c| = 136.8 \pm 5.0$ Å, and $\alpha = \beta = \gamma = 90^\circ$, and is orthorhombic;
 - [5] the crystal according to any one of the above [1] to [4], wherein the crystal has the structural coordinate shown in Figure 4;
- 25 [6] the crystal according to any one of the above [1] to [4], wherein the

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crystal has a structural coordinate different from the structural coordinate as shown in Figure 4 via fluctuation of a protein;

- [7] a three-dimensional structural coordinate of a dipeptidyl peptidase IV, comprising the structural coordinate shown in Figure 4;
- [8] a three-dimensional structural coordinate of a dipeptidyl peptidase IV, comprising a structural coordinate different from the structural coordinate as shown in Figure 4 via fluctuation of a protein;
 - [9] the three-dimensional structural coordinate according to the above [8], wherein the fluctuation of a protein is a state that is caused by molecular oscillation or temperature, and exhibits an activity for the dipeptidyl peptidase IV in a living body;
 - [10] the three-dimensional structural coordinate according to any one of the above [7] to [9], wherein the dipeptidyl peptidase IV is a soluble polypeptide comprising a region located at extramembrane in a full-length human dipeptidyl peptidase IV;
 - [11] the three-dimensional structural coordinate according to any one of the above [7] to [10], wherein the dipeptidyl peptidase IV is a polypeptide having an amino acid sequence in which a transmembrane region is deleted from the amino acid sequence of SEQ ID NO: 2, and a tag peptide is optionally added of to a C-terminal side or N-terminal side thereof:
 - [12] a three-dimensional structural coordinate of a region in a dipeptidyl peptidase IV, comprising the three-dimensional structural coordinate of the region selected from the group consisting of the following (a) to (d):
- (a) a region characterized by Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and

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all or a part of a group of the amino acid residues located in the adjacent area of each of the Ser 630, Asp 708 and His 740 in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate;

- 5 (b) a region characterized by Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and all or a part of a group of the amino acid residues comprising amino acids capable of maintaining physicochemical characteristics physiologically equivalent to each of amino acids in the group of the amino acid residues located in the adjacent area of each of Ser 630, Asp 708 and His 740, in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate,
- (c) a region characterized by a group of amino acid residues comprising amino acids capable of maintaining physicochemical characteristics

 15 physiologically equivalent to each of Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and all or a part of a group of the amino acid residues located in the adjacent area of said group of the amino acid residues in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate; and
 - (d) a region characterized by a group of amino acid residues comprising amino acids capable of maintaining physicochemical characteristics physiologically equivalent to each of Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and all or a part of a group of amino acid residues comprising amino acids

capable of maintaining physicochemical characteristics physiologically equivalent to each of the amino acids in the group of the amino acid residues located in the adjacent area of said group of the amino acids, in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate,

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wherein the region in the dipeptidyl peptidase IV is a region involved in binding or interaction between the dipeptidyl peptidase IV and an effector of the dipeptidyl peptidase IV;

- [13] the three-dimensional coordinate according to the above [12], wherein the physicochemical characteristic is selected from the group consisting of features in shape of a three-dimensional structure, hydrophobicity, electric charge and pK;
 - [14] a method for obtaining a three-dimensional coordinate of a homolog protein of a dipeptidyl peptidase IV, characterized in refining an electron density map of the homolog protein of the dipeptidyl peptidase IV comprising the amino acid sequence of SEQ ID NO: 2, based on all and/or a part of the three-dimensional coordinate of any one of the above [7] to [13], to give a three-dimensional structural coordinate:
- [15] a method for obtaining a three-dimensional structural coordinate of a crystal of a complex of a dipeptidyl peptidase IV and an effector of the dipeptidyl peptidase IV characterized in using all and/or a part of the three-dimensional structural coordinate of any one of the above [7] to [13], to give a three-dimensional structural coordinate;
- [16] a method for identifying pharmacophore of an effector of the dipeptidyl peptidase IV, characterized in identifying the pharmacophore based on all and/or

a part of the three-dimensional structural coordinate of any one of the above [7] to [13], and the steric conformation of the effector of the dipeptidyl peptidase IV; [17] a method for designing, identifying, evaluating or searching an effector of a dipeptidyl peptidase IV, characterized in designing, identifying, evaluating or searching a compound capable of acting on the dipeptidyl peptidase IV, based on all and/or a part of the three-dimensional structural coordinate of any one of the above [7] to [13];

- [18] the method according to the above [17], wherein the method for designing, identifying, evaluating or searching an effector comprises the steps of:
- (i) identifying a region to be targeted for binding or interaction with the effector in a dipeptidyl peptidase IV, based on all and/or a part of the three-dimensional structural coordinate according to any one of the above [7] to [13] and the steric conformation of the effector of the dipeptidyl peptidase IV;
- 15 (ii) identifying atoms or atomic groups capable of generating in the above region at least one intermolecular interaction selected from the group consisting of covalent bond, ionic interaction, ion-dipole interaction, dipole-dipole interaction, hydrogen bonding, van der Waals force, electrostatic interaction and hydrophobic interaction, with the atoms or atomic groups existing in a candidate compound; and
 - (iii) designing a compound based on the information of the above step (i) and/or (ii);
 - [19] the method according to the above [18], wherein the method further comprises the steps of:
- detecting an interaction between the dipeptidyl peptidase IV and the

designed, identified, evaluated or searched candidate compound, wherein when an interaction is detected, the candidate compound is identified as a compound capable of binding to the dipeptidyl peptidase IV, based on a degree of the interaction as an index:

5 the method according to the above [18] or [19], wherein the method further comprises the steps of:

contacting the dipeptidyl peptidase IV with the designed, identified, evaluated or searched candidate compound and measuring the activity of the dipeptidyl peptidase IV.

- wherein when an activity increases or decreases, the designed, identified, 10 evaluated or searched candidate compound is identified as a compound having enhancing action or inhibitory action on the activity of the dipeptidyl peptidase IV, based on a degree of the increase or decrease as an index;
 - an effector of the dipeptidyl peptidase IV obtainable by the method of any one of the above [17] to [20];
 - a program and a medium therefor for use of the three-dimensional structural coordinate of any one of the above [7] to [13], wherein all and/or a part of the three-dimensional structural coordinate of any one of the above [7] to [13] is recorded;
- 20 the program and the medium according to the above [22], comprising a means for identifying, searching, evaluating or designing a compound capable of binding to the dipeptidyl peptidase IV or a compound having an enhancing action or inhibitory action on the activity for the dipeptidyl peptidase IV; and the program and the medium according to the above [23], further [24]
- comprising a means for displaying a three-dimensional graphic display of a 25

molecule.

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BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a photomicrograph of a crystal of a dipeptidyl peptidase IV, wherein the field of view is 4000 $\mu m \times 3000~\mu m$.

Figure 2 is a photograph for X-ray diffraction pattern of a crystal of dipeptidyl peptidase IV.

Figure 3 is a photograph showing a three-dimensional structure of a crystal of a dipeptidyl peptidase IV displayed by the program QUANTA (Accelrys, Inc.).

Figure 4 is a drawing showing a three-dimensional coordinate of a crystal of a dipeptidyl peptidase IV.

BEST MODE FOR CARRYING OUT THE INVENTION

In the present specification, amino acid residues are expressed by using the following abbreviations, which have been adopted by the IUPAC-IUB Commission on Biochemical Nomenclature (CBN). Also, unless explicitly otherwise indicated, the amino acid sequences of peptides and proteins are identified from N-terminal to C-terminal, left terminal to right terminal, the N-terminal being identified as a first residue. Ala: alanine residue; Asp: aspartate residue; Glu: glutamate residue; Phe: phenylalanine residue; Gly: glycine residue; His: histidine residue; Ile: isoleucine residue; Lys: lysine residue; Leu: leucine residue; Met: methionine residue; Asn: asparagine residue; Pro: proline residue; Gln: glutamine residue; Arg: arginine residue; Ser: serine residue; Thr: threonine residue; Val: valine residue; Trp: tryptophane residue;

Tyr: tyrosine residue; Cys: cysteine residue.

The crystal of the present invention is a crystal of a dipeptidyl peptidase IV, having a characteristic sufficient to ensure a resolution capable of analyzing its three-dimensional structure up to the side chain level by X-ray crystallographic structural analysis.

The "characteristic sufficient to ensure a resolution capable of analyzing three-dimensional structure up to the side chain level" is, for example,

- (1) being in a state that a molecule in a unit cell of a crystal has repeats with high regularity, namely, providing diffraction at high resolution;
- 10 (2) having suitable form and size; it is desired that for example, a crystal has at least one side grown to about 0.2 to about 0.5 mm, preferably a cubic crystal having three sides that have similarly grown, or a needle-shaped crystal having a width or thickness of about 0.2 mm or more;
- (3) having chemical stability, dynamic stability and physical stability;
 15 and the like. In a case of the dipeptidyl peptidase IV, which is a polypeptide having a relatively large molecular weight, the term means characteristics sufficient to ensure a resolution of 3Å or less, preferably 2.8Å or less, more preferably 2.6Å or less.

The dipeptidyl peptidase IV used for the preparation of the crystal of the

present invention may have a high purity sufficient for forming the crystal. In
the present invention, the dipeptidyl peptidase IV used for the preparation of the
crystal includes a soluble polypeptide consisting of a region located at
extramembrane in a full-length human dipeptidyl peptidase IV, for example, a
polypeptide in which a transmembrane region in the N-terminal side [namely the
region including the transmembrane sites (the region containing at least the

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amino acid nos: 1-28 of SEQ ID NO: 2, preferably the region of the amino acid nos: 1-32)] is deleted from the amino acid sequence of a full-length human dipeptidyl peptidase IV of SEQ ID NO: 2, and a tag peptide is optionally added to a C-terminal side or N-terminal side of the amino acid sequence. Concrete examples include (I) a polypeptide in which a transmembrane region in the N-terminal side is deleted from the amino acid sequence of a full-length human dipeptidyl peptidase IV of SEQ ID NO: 2; and (II) a polypeptide in which a tag peptide is added to a C-terminal side or N-terminal side of the polypeptide of the above (I). In the polypeptide, since the transmembrane site is deleted therefrom, the polypeptide has excellent characteristics that anchoring to the membrane can be prevented, and the polypeptide is a secretory type and soluble. The tag peptide is not particularly limited. For example, a polyhistidine peptide (an oligopeptide consisting of 4 to 20 of histidine residues) or the like can be preferably used as the tag peptide.

SEQ ID NO: 2 shows the amino acid sequence of a full-length dipeptidyl peptidase IV of human colon.

The full-length dipeptidyl peptidase IV means a polypeptide of a dipeptidyl peptidase IV containing a region comprising a transmembrane site in the N-terminal side. The full-length dipeptidyl peptidase IV includes a polypeptide comprising the amino acid sequence of SEQ ID NO: 2, without being limited thereto, and encompasses its naturally occurring variant, artificially modified variant, a homolog and an ortholog derived from heterogeneous organism, and the like.

Concretely, the full-length dipeptidyl peptidase IV, besides the polypeptide comprising the amino acid sequence of SEQ ID NO: 2, includes

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conservative substitution variants, naturally occurring allelic variants and the like. Also, the full-length dipeptidyl peptidase IV includes a polypeptide having at least one, namely one or more conservative amino acid substitutions, as compared to the polypeptide comprising the amino acid sequence of SEQ ID NO: 2.

The polypeptide as described above may be a polypeptide having biological activities (namely dipeptidyl peptidase IV activity) similar to the polypeptide comprising the amino acid sequence of SEQ ID NO: 2. Concretely, there are included, for instance, a polypeptide having homology of usually about 80% or more, preferably about 90% or more, more preferably about 95% or more on the amino acid level, as compared to the full-length amino acid sequence of SEQ ID NO: 2; a polypeptide encoded by a nucleic acid capable of hybridizing with a nucleic acid consisting of the nucleotide sequence of SEQ ID NO: 1 (nucleotide sequence encoding a full-length dipeptidyl peptidase IV of human colon), under stringent conditions, or a complement thereof; and a polypeptide having deletion, substitution or addition of at least one amino acid, namely one or plural amino acids, preferably one or several amino acids in the amino acid sequence of SEQ ID NO: 2.

The number of deletion, substitution or addition of the amino acids may be to an extent that the biological activities [namely, dipeptidyl peptidase IV activity] are not lost, usually in the number of 1 to about 150, preferably 1 to about 75, more preferably 1 to about 40.

The crystallization is carried out by making a solution containing the desired protein (referred to as a protein solution) supersaturated state, based on the characteristics that the protein in solution state converts to non-soluble state

and precipitates as a crystal when specific conditions are satisfied. Concretely, the protein can be precipitated by the following procedures 1. or 2.:

1. elevating the effective concentration of the protein:

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concretely, adding a precipitant such as a salt, polyethylene glycol or an organic solvent to a protein solution; reducing an amount of a solvent in the protein solution by evaporation or the like; or the like.

2. reducing a repulsive force, or increasing an attractive force between protein molecules:

concretely, adding an organic solvent such as an alcohol to a protein solution; changing a hydrogen ion concentration (pH) or temperature of the protein solution; or the like.

As the conditions for the crystallization, physical and chemical factors such as a hydrogen ion concentration (pH), a kind of buffer used and a concentration thereof, a kind of a precipitant added and a concentration thereof, protein concentration, salt concentration, temperature and the like can be involved. A method for controlling and investigating the factors includes batch methods, dialysis methods, vapor diffusion methods (hanging-drop method, sitting-drop method and the like) and the like, described, for instance, in Blundell, T. L. et al., *PROTEIN CRYSTALLOGRAPHY*, 59-82 (1976), published by Academic Press, or the like.

The method for crystallization includes the batch methods, dialysis methods, vapor diffusion methods and the like. By the above method, physical and chemical factors such as a hydrogen ion concentration (pH), a kind and a concentration of the buffer used, and a kind and a concentration of the precipitant used, and physical and chemical factors such as protein concentration, salt

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concentration and temperature can be also determined.

The hydrogen ion concentration (pH) can be adjusted with a buffer. It is desired that the buffer is a buffer having buffering action in a broad range of pH, and being capable of suppressing precipitation of a non-proteinous crystal between the co-existing ion in the solution used during crystallization and the precipitant or the like. The buffer includes Tris-hydrochloric acid buffer, phosphate buffer, cacodylate buffer, acetate buffer, citrate buffer, glycine buffer and the like.

The precipitant may be a substance capable of elevating an effective concentration of the protein or changing a hydrogen ion concentration (pH) of the protein solution. Generally, the precipitant includes salts such as ammonium sulfate, sodium sulfate, sodium phosphate, potassium phosphate, sodium citrate, ammonium citrate, sodium chloride, potassium chloride and ammonium chloride; polyethylene glycols having various average molecular weights of about 200, about 1000, about 2000, about 4000, about 6000, about 8000, about 20000 or the like; organic solvents such as 2-methyl-2,4-pentadiol, methanol, ethanol, isopropanol, butanol and acetone, and the like.

The protein concentration may be a concentration suitable for crystallization, and it is desired that the protein concentration is, for example, 1 to 50 mg/ml, preferably 5 to 20 mg/ml, more preferably 7 to 15 mg/ml.

It is desired that the temperature conditions are 3° to 25°C, preferably 12° to 22°C.

In the case where the crystallization is carried out by the batch method, the crystallization can be carried out by gradually adding a precipitant solution comprising a precipitant, buffer and the like, so as to form a layer on the top layer of the solution containing the dipeptidyl peptidase IV to give a mixture, or by gradually adding the solution comprising the dipeptidyl peptidase IV, so that the solution is an upper layer of the precipitant solution to give a mixture. Here, the mixture is allowed to stand in a tightly closed vessel.

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In the case where the crystallization is carried out by the dialysis method, the crystallization can be carried out by placing a solution comprising dipeptidyl peptidase IV in a size exclusion semi-permeable membrane, and placing a precipitant solution outside of the size exclusion semi-permeable membrane as a reservoir solution, thereby diffusing the reservoir solution to the solution comprising the dipeptidyl peptidase IV via the semi-permeable membrane.

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In the case where the crystallization is carried out by the hanging-drop method in the vapor diffusion method, the crystallization can be carried out by placing a mixed solution of a solution comprising the dipeptidyl peptidase IV and a precipitant solution in a closed vessel allowing to be hanged at a position above the upper space of a reservoir in which the precipitant solution is contained as a reservoir solution, wherein the vapor pressure of the reservoir solution in the reservoir is set to be lower than that of the mixed solution.

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In the case where the crystallization is carried out by the sitting-drop method in the vapor diffusion method, the crystallization can be carried out by placing a mixed solution comprising a solution comprising the dipeptidyl peptidase IV and a precipitant solution in a closed vessel at a position higher than the liquid surface of a reservoir in which the precipitant solution is contained as a reservoir solution, wherein the vapor pressure of the reservoir solution in the reservoir is set to be lower than that of the mixed solution.

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The crystallization can be carried out by the sitting-drop method from the

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viewpoint of obtaining excellent-quality and large crystal.

When the obtained crystal is a crystal insufficient to ensure the X-ray structural analysis, the crystal may be grown by a seeding method such as macro-seeding method or micro-seeding method, using the obtained crystal as a seed crystal.

When the macro-seeding method is performed, it is desired that the seed crystal is a single crystal that can be isolated by procedures under microscope wherein the seed crystal has excellent external form (having excellent crystallinity). Also, it is desired that the seed crystal is washed with a drop of a solution obtained by diluting the precipitant, for example, by 0.5 to 1.0-fold. It is desired that the solution used for seeding of the seed crystal is a protein solution having a degree of supersaturation that the crystal grows but the crystal nuclei do not grow. On the other hand, when the micro-seeding method is performed, the form and size of the seed crystal are not particularly limited.

The sequence information for the dipeptidyl peptidase IV and cDNA encoding the dipeptidyl peptidase IV can be obtained from a known information source [GenBank/EMBL accession No: X60708; Misumi et al., *Biochim. Biophys. Acta*, 1131, 333-336, (1992); GenBank/EMBL accession No: M80536; Darmoul et al., *J. Biol. Chem.*, 267, 4824-4833, (1992)]. Therefore, the dipeptidyl peptidase IV or a soluble polypeptide thereof can be produced by using conventional means for gene engineering on the basis of the above sequence information.

The nucleic acid used for production of the dipeptidyl peptidase IV or a soluble polypeptide thereof may be any nucleic acid in which the encoded polypeptide exhibits a dipeptidyl peptidase IV activity. For example, a nucleic

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acid encoding a polypeptide consisting of the amino acid sequence in which a transmembrane region in the N-terminal side (a region containing at least the amino acid nos: 1-28, preferably the region of the amino acid nos: 1-32) is deleted from the full-length human dipeptidyl peptidase IV, and a tag peptide is optionally added to a C-terminal side or N-terminal side of the amino acid sequence.

The nucleic acid can be obtained by, for instance, obtaining a fragment comprising a nucleic acid encoding a full-length dipeptidyl peptidase IV or a part thereof by means of conventional DNA recombination technique, and appropriately arranging the obtained fragment.

SEQ ID NO: 1 shows a sequence of a nucleic acid encoding a full-length dipeptidyl peptidase IV of human colon.

The nucleic acid (DNA or RNA) encoding a full-length dipeptidyl peptidase IV includes, for instance, a nucleic acid comprising human nucleic acids comprising the nucleotide sequence of SEQ ID NO: 1 without being limited thereto, and includes its naturally occurring variant, artificially modified variant, a homolog or ortholog derived from heterogeneous organism.

In other words, besides the nucleic acid comprising the nucleotide sequence of SEQ ID NO: 1, the nucleic acid includes a nucleic acid capable of hybridizing with a nucleic acid comprising the nucleotide sequence of SEQ ID NO: 1 under stringent conditions, more preferably under high-stringent conditions), or a complement thereof (nucleic acid having a complementary sequence).

Concrete examples of the nucleic acid described above include, for instance, a nucleic acid having usually about 70% or more, preferably about 80%

or more, more preferably about 85% or more, still more preferably about 90% or more, still more preferably about 95% or more, homology to the nucleotide sequence of SEQ ID NO: 1, and it is preferable that the polypeptide encoded by the above nucleic acid has a dipeptidyl peptidase IV activity.

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The dipeptidyl peptidase IV activity can be measured by, for example, incubating in a 1.5 ml reaction mixture [composition: 1.5 mM substrate (Gly-Pro-paranitroanilide), 71 mM glycine-NaOH (pH 8.7)] at 37°C for 10 minutes, and determining the liberated paranitroanilide at the absorbance of 405 nm. One unit (1 U) of a dipeptidyl peptidase IV is defined as an amount of the enzyme required for liberating 1 µmol of paranitroanilide per 1 minute.

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In the present invention, the hybridization under stringent conditions can be carried out as normal stringent conditions by performing hybridization in a hybridization solution having a salt concentration of $6 \times SSC$ or an equivalent concentration thereto, under the temperature conditions of 50° to $70^{\circ}C$ for about 16 hours, and optionally performing pre-washing with a solution having a salt concentration of $6 \times SSC$ or an equivalent concentration thereto, and thereafter performing washing with a solution having a salt concentration of $1 \times SSC$ or an equivalent concentration thereof. Furthermore, as the conditions having still higher stringency (high-stringent conditions), the hybridization can be carried out by washing with a solution having a salt concentration of $0.1 \times SSC$ or an equivalent concentration thereto in the above method.

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The dipeptidyl peptidase IV used for the crystallization has purity that can form a crystal, and the purity can be confirmed by conventional means of confirming purity (for example, a method comprising electrophoresing a fraction by polyacrylamide gel electrophoresis, SDS-polyacrylamide gel electrophoresis

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or the like, and visualizing the fraction by silver staining, or the like).

The X-ray structural analysis data of the crystal can be obtained by subjecting the crystal of the present invention to an X-ray crystallographic structural analysis known to one of ordinary skill in the art [for example, see Blundell, T. L. et al., PROTEIN CRYSTALLOGRAPHY, 59-82 (1976), published by Academic Press, and the like], whereby a three-dimensional structural coordinate (a value showing the relationship of the spatial positions of each atom) and a three-dimensional structure model for the crystal can be obtained. Concretely, the three-dimensional structural coordinate of the dipeptidyl peptidase IV is obtained as an atomic coordinate by procedures comprising the steps of 1) irradiating the crystal of the present invention with a monochromatic X-ray to give an X-ray diffraction pattern, 2) obtaining X-ray diffraction intensity data from the X-ray diffraction pattern, 3) obtaining an electron density map by Fourier transform, and 4) allocating a polypeptide chain and side chain thereof on the electron density map based on the amino acid sequence of the polypeptide used for the crystal. Furthermore, the three-dimensional structure is clarified by molecule-modeling based on the three-dimensional structural coordinate. Therefore, the three-dimensional structural coordinate of the dipeptidyl peptidase IV obtained from the crystal of the present invention is also encompassed within the scope of the present invention.

The crystallographic parameters for the crystal are obtained from the X-ray diffraction intensity data of the crystal of the present invention. The crystal of the present invention is an orthorhombic crystal having a space group of $P2_12_12_1$, and a lattice constant of the unit cell of $|a| = 118.0 \pm 5.0$ Å, $|b| = 125.9 \pm 5.0$ Å, $|c| = 136.8 \pm 5.0$ Å, and $\alpha = \beta = \gamma = 90^{\circ}$. The crystal has a

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2.6Å resolution by X-ray crystallographic structural analysis, that is, the crystal has characteristics sufficient to ensure a resolution capable of analyzing up to the side chain level of the polypeptide.

It is a known fact to one of ordinary skill in the art that the same protein can be crystallized even under different conditions. Therefore, the present invention is not limited to only the conditions for crystallization, and the crystal that shows substantially the same crystallographic constants as those in the present invention are also encompassed within the scope of the present invention.

More concretely, the crystal of the dipeptidyl peptidase IV of the present invention has a structural coordinate as shown in Figure 4, or a structural coordinate different from the structural coordinate as shown in Figure 4 via fluctuation of a protein.

The crystal according to the present invention can also be used as a seed crystal for carrying out the crystallization of a polypeptide having a three-dimensional structure similar to that of the dipeptidyl peptidase IV used for, for example, carrying out the crystallization of the dipeptidyl peptidase IV, dipeptidyl peptidase IV-like proteins, homolog proteins and the like, which are derived from other organism species.

When the crystal of the present invention is irradiated with X-ray, a low-temperature measurement may be carried out, as described in Examples set forth below.

The X-ray structural analysis data are converted to a structure factor by evaluating the intensity of X-ray diffraction using MOSFILM Program Package (Version 6.1). Also, in order to obtain the information for the phase, multiple isomorphous replacement method or the like can be performed, for example, as

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described in Examples.

In the structural analysis, CCP4 (Collaborative Computational Project, Number 4, 1994, "The CCP4 Suite: Programs for Protein Crystallography," Acta Cryst. D50, 760-763) program or the like is used.

The three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention can be obtained, for example, as follows. Firstly, Fourier transform calculation is carried out using the differences between the diffraction intensity obtained from two kinds of isomorphous replacement crystals of mercury and the diffraction intensity obtained from native crystal, and investigating the large peaks provided by the heavy atoms (mercury) on the Patterson's diagram to determine the locations of each mercury atoms in the unit cell of the real space. The phase of the crystal structure factor for the native crystal is determined using the obtained location coordinate for the mercury atoms. Furthermore, refinement is performed using the crystal structure factor of the native crystal and two kinds of the crystal structure factors of the isomorphous replacement crystals of mercury, and the coordinate for each of the mercury atoms is more accurately determined. An electron density map for the crystal of the dipeptidyl peptidase IV in the real space is obtained using the phase of the crystal structure factor of the native crystal calculated from the refined mercury atoms coordinate. Furthermore, the electron density map is improved by performing smoothing and histogram matching for the electron density map of the solvent region, whereby an electron density map necessary and sufficient for building a molecular model can be obtained. Next, the sites corresponding to the amino acid residues of the dipeptidyl peptidase IV on the electron density map are identified using QUANTA (manufactured by Accelrys, Inc.) to build the

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molecular model to give a three-dimensional structural coordinate.

The three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention is shown in Figure 4. Figure 4 shows the obtained three-dimensional structural coordinates, according to the format of the Protein Data Bank, which is a notation generally used by one of ordinary skill in the art.

The three-dimensional structural coordinates shown in Figure 4 are those represented using the origin of the unit cell of the crystal as the origin of the three-dimensional space. The R factor that is considered as an index for the accuracy of the obtained molecular model is 24.89%, and the free R factor is 30.15%. In addition, the deviation in the interatomic bond distance from the ideal state of the three-dimensional structure (rms-deviation) and the deviation in the bond angle are 0.006Å and 1.305°, respectively. In the case, for instance, the three-dimensional structural coordinate of the present invention is used for the calculation by a computer, a novel structural coordinate obtained as a result of the operation for mathematical transfer, such as translation, rotation, or symmetry in the three-dimensional space without changing the relative configuration of the atoms, is also encompassed within the scope of the present invention. Furthermore, not only all of the three-dimensional structural coordinate of the present invention but also a part thereof are also encompassed within the scope of the present invention.

The three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention can be used, for example, as shown in Figure 3, for three-dimensional graphic displaying of the stereogram of the three-dimensional structure model, and for evaluation of the structure-activity relationship and the quantitative structure-activity relationship. Also, the structural features of the

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crystal of the present invention can be more concretely shown using the three-dimensional structural coordinate shown in Figure 4. The evaluation of the structure-activity relationship or quantitative structure-activity relationship by the three-dimensional structure model is also encompassed within the scope of the present invention.

According to the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, one of the characteristics of the dipeptidyl peptidase IV can be found in that the dipeptidyl peptidase IV has 273 molecules of bond water in an asymmetric unit and has 5 molecules of N-acetylglucosamine residues per 1 molecule of the dipeptidyl peptidase IV.

According to the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, the information for atoms or atomic groups of the side chain of the dipeptidyl peptidase IV, interacting with the atoms or atomic groups of a known effector of the dipeptidyl peptidase IV via an intermolecular interaction can be obtained.

Furthermore, according to the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, the information of regions in the dipeptidyl peptidase IV that are susceptible to binding or intermolecular interaction with the effector can be obtained.

In addition, according to the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, the information of the structure specific to the dipeptidyl peptidase IV, which is not found in proteins other than the dipeptidyl peptidase IV, can be obtained. Therefore, higher selectivity in the effector targeting a protein other than the dipeptidyl peptidase IV can be designed, when the effector also acts on the dipeptidyl peptidase IV.

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The intermolecular interaction includes covalent bond, ionic interaction, ion-dipole interaction, dipole-dipole interaction, hydrogen bonding, van der Waals force, electrostatic interaction, hydrophobic interaction and the like.

In the present specification, the atoms or atomic groups of the effector and atoms or atomic groups of the side chain of the dipeptidyl peptidase IV, which interact with each other via intermolecular interaction, are referred to as "pharmacophore."

Also, according to the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, the information for the structure specific for the dipeptidyl peptidase IV, which is not found in proteins other than the dipeptidyl peptidase IV, can be provided.

In addition, for example, when the measurement conditions are different in X-ray diffraction, or the three-dimensional structure of the complex in the solution is analyzed using multidimensional NMR, and the like, the three-dimensional structural coordinate may differ from that shown in Figure 4. The three-dimensional structural coordinate varies depending on the fluctuation of protein and the like, and is encompassed within the scope of the present invention.

In the present specification, the "fluctuation of protein" means a state that is caused by molecular oscillation, temperature and the like, and accompanied with the structural change that can exhibit an activity for the dipeptidyl peptidase IV in a living body.

Also, according to the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, one of the characteristics of the dipeptidyl peptidase IV resides in that the amino acid residues, Ser 630, Asp 708

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(b)

and His 740, which are involved in the activity deduced by experiments by using various active inhibitors of the dipeptidyl peptidase IV, exist in the adjacent area, even though the amino acid residues exist in distant locations on the primary sequence. Concretely, the distance between the $O_{\delta 2}$ atom of Asp 708 and the $N_{\delta 1}$ atom of His 740, and the distance between the $N_{\epsilon 2}$ atom of His 740 and the O_{γ} atom of Ser 630 are distances that can form hydrogen bonding.

Therefore, the present invention also includes a three-dimensional structural coordinate of the region in the dipeptidyl peptidase IV, which is involved in binding or interaction of the dipeptidyl peptidase IV with an effector thereof, including a three-dimensional structural coordinate of a region selected from the group consisting of the following (a) to (d):

- a region characterized by Ser 630, Asp 708 and His 740 in the amino acid (a) sequence of SEQ ID NO: 2, and all or a part of a group of amino acid residues located in the adjacent area of each of the Ser 630, Asp 708 and His 740 in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate;
- a region characterized by Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and all or a part of a group of amino acid residues comprising amino acids 20 capable of maintaining physicochemical characteristics physiologically equivalent to each of amino acids of the group of amino acid residues located in the adjacent area of each of Ser 630, Asp 708 and His 740, in the structural coordinate shown in Figure 4 or the three-dimensional 25 structure model defined by the structural coordinate.

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- aregion characterized by a group of amino acid residues comprising amino acids capable of maintaining physicochemical characteristics physiologically equivalent to each of Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and
- all or a part of a group of amino acid residues located in the adjacent area of said group of the amino acid residue in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate; and
- (d) a region characterized by a group of amino acid residues comprising
 amino acids capable of maintaining physicochemical characteristics
 physiologically equivalent to each of Ser 630, Asp 708 and His 740 in the
 amino acid sequence of SEQ ID NO: 2, and

all or a part of a group of amino acid residues of a group of amino acid residues comprising amino acids capable of maintaining physicochemical characteristics physiologically equivalent to the each amino acid of the amino acid residues located in the adjacent area of said groups of the amino acids, in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate.

In the present specification, the "adjacent (area)" refers to an area involved in covalent bond, ionic interaction, ion-dipole interaction, dipole-dipole interaction, hydrogen bonding, van der Waals force, electrostatic interaction, hydrophobic interaction or the like with the amino acid residues, concretely, a region within 10Å, preferably within 8Å, more preferably within 5Å.

The physicochemical characteristic includes features in shape of the three-dimensional structure, hydrophobicity, electric charge, pK and the like.

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The "amino acid capable of maintaining physicochemical characteristics physiologically equivalent" may be an amino acid analogue residue obtained by replacing a side chain of amino acid residues in the three-dimensional structural coordinate shown in Figure 4 with other side chain, for example, showing bioisosterism. Alternatively, the amino acid residue in the three-dimensional structural coordinate shown in Figure 4, may be replaced with another amino acid residue belonging to the same Group, in any of the following Groups I to VI:

I glycine, alanine;

10 II valine, isoleucine, leucine;

III aspartic acid, glutamic acid, asparagine, glutamine;

IV serine, threonine;

V lysine, arginine;

VI phenylalanine, tyrosine.

According to the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, a three-dimensional structural coordinate of a polypeptide can be easily derived if an accurate amino acid sequence is determined, even when the polypeptide is a dipeptidyl peptidase IV or a dipeptidyl peptidase IV-like protein derived from other organism species, as long as the polypeptide is a polypeptide having high homology on the level of amino acid sequence with the dipeptidyl peptidase IV used for the preparation of the crystal of the present invention (for example, at least 20%, preferably 30% or more, more preferably 40% or more).

Furthermore, the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention can be used for X-ray crystallographic

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structural analysis of the crystal and the like of other proteins having an amino acid sequence with significant homology with the dipeptidyl peptidase IV used for the preparation of the crystal of the present invention. Concretely, according to the molecular replacement method [for example, see Blundell, T. L. et al., PROTEIN CRYSTALLOGRAPHY, 446-464 (1976), published by Academic Press and the like], the three-dimensional structural coordinate thereof can be quickly and readily obtained from the structure factors obtained by the X-ray diffraction pattern of the crystal, without using multiple isomorphous replacement method, even for the determination of the structural coordinate of the above-mentioned crystal of which structural coordinate has not yet been known.

In the present specification, the term "significant homology" is a case where there is identity of 20%, or more, preferably by 30% or more, between the amino acid sequences.

When the molecular replacement method is performed, for example, a program such as X-PLOR and CNX (both manufactured by Accelrys Inc.) or AMORE [one of the programs of CCP4 (Collaborative Computational Project, Number 4), *Acta Crystallogr.* **D50**, 670-673 (1994)] can be run by a computer on which the program can be executed. Here, whether or not the molecular replacement method is applicable can be determined by actually applying the molecular replacement method to the structure factors calculated from the X-ray diffraction pattern of the desired crystal and obtaining a significant solution.

In other words, the three-dimensional structural coordinate obtained by structural analysis by molecular replacement method is encompassed within the scope of the present invention as long as a significant solution is obtained. The present invention also encompasses a three-dimensional structural coordinate of

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a dipeptidyl peptidase IV, or a dipeptidyl peptidase IV-like protein, namely a homolog protein or the like of other organism species derived by the above method.

Therefore, according to the present invention, a method for obtaining a three-dimensional structural coordinate of a homolog protein of a dipeptidyl peptidase IV comprising the step of performing refinement of an electron density map of the homolog protein of the dipeptidyl peptidase IV comprising the amino acid sequence of SEQ ID NO: 2, based on the three-dimensional structural coordinate of the present invention, to give a three-dimensional structural coordinate is provided. Also, a method for obtaining a three-dimensional structural coordinate of a crystal of a complex of a dipeptidyl peptidase IV and an effector of the dipeptidyl peptidase IV, based on the three-dimensional structural coordinate of the present invention, is likewise provided.

According to the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, a method for identifying a region or site for a target for binding or interaction between the dipeptidyl peptidase IV and an effector of the dipeptidyl peptidase IV is provided, based on the analysis of the binding regions between the dipeptidyl peptidase IV and a known effector of the dipeptidyl peptidase IV such as an inhibitor, or based on the simulation of the interaction between the dipeptidyl peptidase IV and a known effector of the dipeptidyl peptidase IV.

Also, based on the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention and the steric conformation of the effector of the dipeptidyl peptidase IV, the pharmacophore of the effector of the dipeptidyl peptidase IV can be identified. A method for identifying the

pharmacophore is also provided. The method is useful for designing an effector having excellent characteristics such as higher avidity, higher biological activity, higher biological stability, higher thermodynamic stability, higher absorbency to a living body, and lower toxicity.

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Concretely, for example, the region or site for a target involved in binding or interaction of the dipeptidyl peptidase IV and an effector of the dipeptidyl peptidase IV, can be identified by:

- 1) obtaining a crystal of a complex of the dipeptidyl peptidase IV and a known effector of the dipeptidyl peptidase IV such as an inhibitor, and obtaining a three-dimensional structural coordinate of the crystal based on the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention and the steric conformation of the effector of the dipeptidyl peptidase IV, whereby obtaining the three-dimensional structural coordinate of a binding region of the dipeptidyl peptidase IV and the effector:
- 15 2) simulating an intermolecular interaction between the dipeptidyl peptidase IV and a known effector of the dipeptidyl peptidase IV based on the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention and the steric conformation of the effector of the dipeptidyl peptidase IV;

or the like.

The crystal of the above-mentioned complex can be obtained by, for example, incubating the crystal of the present invention in a solution comprising the effector, forming a complex of the dipeptidyl peptidase IV and the effector, and crystallizing the obtained complex, and the like.

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Also, when the three-dimensional structural coordinate of the crystal of

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the complex is obtained, the steric structure of the effector of the abovementioned complex can be readily obtained by calculating the differential Fourier diagram utilizing a three-dimensional structure model defined by the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, whereby specific interaction forms and interaction sites between the dipeptidyl peptidase IV and the effector can be readily clarified.

When the intermolecular interaction is simulated, for example, the space regions, residues and the like in which covalent bond, ionic interaction, ion-dipole interaction, dipole-dipole interaction, hydrogen bonding, van der Waals force, electrostatic interaction, hydrophobic interaction or the like can be simulated, based on the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention and the steric conformation of the effector of the dipeptidyl peptidase IV.

Furthermore, according to the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, the three-dimensional structural coordinate or the three-dimensional structure model based on the three-dimensional structural coordinate regarded as an active center of the dipeptidyl peptidase IV, sites indirectly acting on the active center and regions or sites involved in binding or interaction with the effector, or the like, is obtained, whereby a compound capable of specifically acting on the dipeptidyl peptidase IV can be designed, identified, evaluated or searched.

For example, in the structural coordinate of Figure 4 and the threedimensional structure model defined by the structural coordinate, a compound capable of modifying the activity of the dipeptidyl peptidase IV can be designed, identified, evaluated or searched, based on the regions characterized by Ser 630,

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Asp 708 and His 740, and all or a part of amino acid residues of the group of the amino acid residues located in the adjacent area of the Ser 630, Asp 708 and His 740.

Therefore, according to the present invention, a method for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV is provided.

One of the significant features of the method of the present invention for designing, identifying, evaluating or searching an effector resides in that the method comprises designing, identifying, evaluating or searching a compound capable of acting on the dipeptidyl peptidase IV, based on the three-dimensional structural coordinate of the present invention.

According to the method of the present invention for designing, identifying, evaluating or searching an effector, since the method is based on the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, the information for a structure specific to the dipeptidyl peptidase IV, which is not found in proteins other than the dipeptidyl peptidase IV can be obtained. Therefore, according to the method of the present invention for designing, identifying, evaluating or searching an effector, the method has an excellent effect that the selectivity of the effector of the dipeptidyl peptidase IV can be enhanced.

Also, according to the method of the present invention for designing, identifying, evaluating or searching an effector, since the method is based on the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention, visual studies and/or energy calculation can be made according to the method by using a computer and the like. Therefore, there are

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exhibited some excellent effects that a compound having excellent characteristics such as having higher avidity, higher biological activity, higher biological stability, higher thermodynamic stability, higher absorbency in a living body, and lower toxicity, than those for a known inhibitor can be designed, identified, evaluated or searched, and that logical design can be performed in the three-dimensional space.

In the present specification, the "effector" includes a compound that inhibits or enhances the activity (i.e. inhibitor or activator), which may be natural compounds or synthetic compounds, or may be polymers or low-molecular weight compounds.

A concrete example of the method of the present invention for designing, identifying, evaluating or searching an effector includes a method comprising the steps of:

- (i) identifying a region to be targeted for binding or interaction with the effector in a dipeptidyl peptidase IV, based on all and/or a part of the three-dimensional structural coordinate of the present invention and the steric conformation of the effector of the dipeptidyl peptidase IV;
- (ii) identifying corresponding atoms or atomic groups capable of generating in the region at least one intermolecular interaction selected from the group consisting of covalent bond, ionic interaction, ion-dipole interaction, dipole-dipole interaction, hydrogen bonding, van der Waals force, electrostatic interaction and hydrophobic interaction, with the atoms or atomic groups existing in a candidate compound; and
- (iii) designing a compound based on the above information of the above step(i) and/or (ii).

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The three-dimensional structural coordinate used for designing, identifying, evaluating or searching a compound capable of binding to the dipeptidyl peptidase IV may be a coordinate fixed in the three-dimensional space, and the intensity of binding with the compound or the like can be calculated by carrying out translation or rotation in the three-dimensional space, and transfer to an extent that the chemical covalent bond would not be cleaved in the amino acid residues of the dipeptidyl peptidase IV.

In the above step (i), the "region to be targeted in the dipeptidyl peptidase IV" preferably includes an active center of the dipeptidyl peptidase IV, sites indirectly acting on the active center and the like. For example, there is included a region characterized by Ser 630, Asp 708 and His 740 and all or a part of a group of the amino acid residues located in the adjacent area of Ser 630, Asp 708 and His 740, and the like in the structural coordinate of Figure 4 and the three-dimensional structure model defined by the structural coordinate. The atoms or atomic groups that can be matched therewith are identified, based on the three-dimensional structural coordinate of an active center, sites indirectly acting on the active center and the like, whereby the candidate atoms or candidate atomic groups can be obtained.

In the above step (ii), for example, the atoms or atomic groups capable of associating via intermolecular interaction between the atoms or atomic groups in the region, concretely, the corresponding atoms or atomic groups capable of generating covalent bond, ionic interaction, ion-dipole interaction, dipole-dipole interaction, hydrogen bonding, van der Waals force, electrostatic interaction, hydrophobic interaction and the like, are searched and extracted, based on the information identified in the above step (i).

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Next, in the above step (iii), the corresponding atoms or atomic groups searched in the above step (i) and/or (ii) are combined to design a compound.

Thereafter, if desired, whether or not the compound designed in the above step (iii) is matched via intermolecular interaction with the side chains and atoms or atomic groups in the dipeptidyl peptidase IV as defined by the three-dimensional structural coordinate of the present invention can be simulated.

The compound designed, identified, evaluated or searched by the above steps (hereinafter also referred to as a candidate compound in the present specification) can be obtained by generally used chemical synthetic methods, depending on the compound.

In addition, in the method of the present invention for designing, identifying, evaluating or searching an effector, there can be carried out a step of detecting the interaction between the dipeptidyl peptidase IV and the candidate compound. When the interaction is detected, the interaction serves as an index showing that the above candidate compound is a compound capable of binding to the dipeptidyl peptidase IV.

The above interaction can be detected by, for example, plasmon resonance analysis apparatus, mass spectrometer, titration isothermal calorimetry, NMR and the like. For example, in the case of plasmon resonance analysis apparatus, when a sensorgram indicates the formation of a complex, by contacting the dipeptidyl peptidase IV-immobilized matrix with the candidate compound and performing analysis by optical detection (for example, photometer, polarization photometer and the like) and the like, it would be an index showing that the interaction between the candidate compound and the dipeptidyl peptidase IV is generated. For example, in the case of a mass spectrometer, when a spectrum

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indicates the formation of a complex, by contacting the dipeptidyl peptidase IVimmobilized matrix with the candidate compound and performing analysis with a mass spectrometer (matrix-assisted laser desorption/ionization-time of flight mass spectrometry: MALDI-TOF MS, electro spray-ionization mass spectrometer: ESI-MS and the like), it would be an index showing that the interaction between the candidate compound and the dipeptidyl peptidase IV is generated. For example, in the case of titration-thermal calorimetry interaction analysis, when the titration curve indicates the formation of a complex, by contacting a solution of the dipeptidyl peptidase IV with the candidate compound, and measuring the heat coming in and out of a thermal diode and the like, it would be an index showing that the interaction between the candidate compound and dipeptidyl peptidase IV is generated. For example, in the case of NMR, when a spectrum indicates the formation of a complex, by analyzing by NMR a solution prepared mixing the dipeptidyl peptidase IV and a candidate compound, it would be an index showing that the interaction between the candidate compound and the dipeptidyl peptidase IV is generated.

Furthermore, the method of the present invention for designing, identifying, evaluating or searching an effector may further comprise the steps of contacting the dipeptidyl peptidase IV with a candidate compound, and thereafter measuring the activity of the dipeptidyl peptidase IV. When the dipeptidyl peptidase IV activity increases or decreases, it would be an index showing that the candidate compound is a compound having enhancing action or inhibitory action on the activity of the dipeptidyl peptidase IV.

The dipeptidyl peptidase IV activity can be measured by, for example, incubating a 1.5 ml reaction mixture [composition: 1.5 mM substrate

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(Gly-Pro-paranitroanilide), 71 mM glycine-NaOH (pH 8.7)] at 37°C for 10 minutes in the presence of a candidate compound, and measuring the liberated paranitroanilide at the absorbance of 405 nm. During the measurement of the activity, the candidate compound may be evaluated by using a reaction system in which a suitable dilution series of the compound is added thereto.

The method of the present invention for designing, identifying, evaluating or searching the effector can be performed by, for example, sequentially selecting the interaction between the dipeptidyl peptidase IV and the compounds in a database in a computer to which the structures of plural of compounds had been inputted, or the interaction between the dipeptidyl peptidase IV and the designed compound, by visual methods (visual selection method) utilizing the database; and/or sequentially calculating the avidity with a computer, and searching a compound capable of stably interacting with the dipeptidyl peptidase IV from the database (computer-assisted avidity evaluation method) and the like, based on the three-dimensional structural coordinate of the present invention.

In the above visual selection method, the database of the structures of compounds may be a database in which the three-dimensional structural coordinates have been determined and inputted. Alternatively, in the case of a compound having a low molecular weight, the database may be a database in which the information for chemical covalent bond of a compound having a low molecular weight had been inputted, because the conformation can be relatively freely changed, and the three-dimensional structural coordinate of each conformation can be derived by calculation in a relatively short time.

Concretely, in the visual selection method, the expected complex between the dipeptidyl peptidase IV and a candidate compound or a part thereof is firstly

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displayed on a computer screen, based on the three-dimensional structural coordinate of the present invention. Next, the intermolecular interaction binding between a compound in the database and the binding regions of the dipeptidyl peptidase IV is simulated on the computer, taking chemical interaction into consideration. Also, the simulation of the chemical modification of the compound is performed on the computer, and the changes in the interaction caused as a result thereof are observed on the computer screen. During the simulation, the three-dimensional space can be more easily understood by displaying the three-dimensional structure of the protein on the computer screen so that the structure corresponds to Crystal Eye glasses supplied by Silicone Graphics; simultaneously displaying two screens in which each angle is adjusted for displaying the object, according to the visual fields of the right eye and left eye, which is so-called referred to as "stereovision" which is frequently used by one of ordinary skill in the art; or the like. In addition, the three-dimensional structure can be visually studied by methods other than the stereoscopic displaying of the three-dimensional structure.

The candidate compound capable of generating suitable interaction can be obtained by displaying on a computer a group of candidates with appropriate conformation and selecting an appropriate one therefrom; calculating a structure having a low energy state on a computer; or the like. Next, a derivative of a compound capable of generating more preferable binding with the dipeptidyl peptidase IV may be searched among the candidate compound.

More specifically, on the level of the three-dimensional structure, the followings may be taken into consideration:

25 a group likely to be charged negatively, such as carboxyl group, nitro

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group, or a halogen group in the compound interacts with an amino acid residue having a positive charge, such as lysine, arginine or histidine in the dipeptidyl peptidase IV;

- a group likely to be charged positively, such as amino group, imino group or guanidyl group in the compound interacts with an amino acid residue having negative charge, such as glutamic acid or aspartic acid in the dipeptidyl peptidase IV;
- a hydrophobic functional group such as an aliphatic group or an aromatic group in the compound interacts with a hydrophobic amino acid residue such as alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophane or methionine in the dipeptidyl peptidase IV;
 - a group involved in hydrogen bonding, such as hydroxyl group or amide group is allowed to form hydrogen bonding with a main chain or side chain portion;
- a group or an atom likely to be charged negatively, such as carboxyl group, nitro group or a halogen group in the compound interacts with a positively charged atom on a main chain or side chain portion;
 - a group or an atom likely to be charged positively, such as amino group, imino group or guanidyl group in the compound interacts with a negative charged atom on a main chain or a side chain portion;
 - the flexibility of the three-dimensional structure of the compound is lowered by, for instance, cyclizing the linear chain portion; or the like. For example, a derivative may be designed and synthesized so that the atoms having negative charge of the candidate compound are located in the adjacent region of the side chain of an amino acid residue having positive charge

such as lysine, arginine or histidine, in the amino acid residue of the dipeptidyl peptidase IV, and that an atom having positive charge of the candidate compound is located in the adjacent region of the side chain of the amino acid residue having negative charge such as glutamic acid or aspartic acid in the amino acid residue of the dipeptidyl peptidase IV. Also, a group suitable for forming a hydrophobic interaction may be introduced into the portion capable of forming a hydrophobic interaction between the compound and the dipeptidyl peptidase IV, to design and synthesize a derivative. In addition, a group suitable for forming hydrogen bonding may be introduced into the portion capable of forming hydrogen bonding between the compound and the dipeptidyl peptidase IV, to design and synthesize a derivative. In the above-mentioned designing, it is desirable that van der Waals interaction is as high as possible, and that steric hindrance does not occur between each of the atoms. Furthermore, it is desirable that new void portions are not produced by modification of the compound and that in regions already containing void portions, the void portions are filled as much as possible.

As described above, the design, identification, evaluation or searching of a final compound can be thus performed with visually comprehensively considering intermolecular interaction and other factors on a computer screen.

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In the computer-assisted avidity evaluation method, in order to determine the validity for the designing of a new compound, and to obtain a compound that can stably interact from the compounds in the database, a docking software (DOCK, GOLD, FlexX, Glide or the like) is used for evaluation of binding based on the energy by calculating a molecular force field between the compound and the dipeptidyl peptidase IV, evaluation of binding based on chemical

characteristics, evaluation of binding based on the Protein Data Bank (PDB), and the like. Further, in a model system consisting of the compound and the dipeptidyl peptidase IV, or in a model system further comprising solvent molecules and the like, it can be led to a compound that can stably interact by obtaining the index showing avidity, such as free energy of bonding, the ratio obtained from bond state number and non-bond state number, and the like by using molecular kinetic calculation or Monte Carlo calculation. The programs for calculation of molecular force field and molecular kinetic include AMBER, CHARMm, DISCOVER, PRESTO and the like, and the force field used includes AMBER, CHARMm, OPLS, MMCF, CVFF and the like. Furthermore, a program such as Ludi which automatically outputs the candidates for a candidate compound by providing a three-dimensional structural coordinate of the amino acid residues interacting in the dipeptidyl peptidase IV may be used.

The visual selection method and computer-assisted avidity evaluation method can be performed alone or in combination. In the case of performing the methods in combination, the avidity is actually calculated for the compounds that has been expected to be more desirable in visual investigation, and the validity thereof is evaluated. By repeatedly performing the calculation and evaluation, more excellent compounds may be designed, identified, evaluated or searched.

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Next, the designed, identified, evaluated or searched compound is optimized to be a more excellent compound, such as a compound having more excellent characteristics as a medicament, such as being excellent *in vivo* kinetics, having low toxicity and low side-effect; a compound having a still higher biological activity as an effector; a compound having an advantageous structure as a medicament in view of its oral administration; and the like.

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logically.

The resulting candidate compound can be obtained using generally used techniques for chemical synthesis depending on the kind of the compound.

The present invention also encompasses an effector of the dipeptidyl peptidase IV, which is obtained by the method of the present invention for designing, identifying, evaluating or searching an effector. When the effector is a compound capable of inhibiting or enhancing the activity of the dipeptidyl peptidase IV, the effector (inhibitor or activator) is expected to be an agent for, for example, a modulatory agent of immune response, a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like.

The three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention can be provided as a computer program, a medium or the like, which displays the three-dimensional structure of the molecule based on the three-dimensional structural coordinate and can be provided via a telecommunication line or the like. Therefore, using a computer or the like, the three-dimensional coordinate of the dipeptidyl peptidase IV can be displayed in detail, allowing to perform the method of the present invention for designing, identifying, evaluating or searching an effector more rapidly, conveniently and

The present invention also encompasses a program or a medium therefor for use of the three-dimensional structural coordinate, in which all and/or a part of the three-dimensional structural coordinate of the dipeptidyl peptidase IV of the present invention is recorded.

The medium may be any of those in which the three-dimensional structural coordinate of the present invention can be derived on a program that

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runs on a computer, and includes, for instance, electric memory media referred to as memory; semi-permanent memory media such as a FD, a hard disk, an optical disk, an opto-magnetic disk and a magnetic tape, and the like. In addition, the program and the medium therefor for use of the three-dimensional structural coordinate of the present invention also encompass those having a form which can be communicated via a telecommunication line such as internet.

Also, the program and the medium therefor for use of the three-dimensional structural coordinate of the present invention may further comprise a means for displaying the three-dimensional graphic display of the molecule. The program or the medium therefor which comprises the means for displaying the three-dimensional graphic display has advantages that visual studies and/or calculation of avidity can be made more conveniently, so that there is more facilitated a logical design on the three-dimensional structural level for obtaining a compound having excellent characteristics such as higher avidity, higher biological activity, higher biological stability, higher thermomechanical stability, higher absorbency to a living body, and lower toxicity than those for known effectors of the dipeptidyl peptidase IV.

As the means capable of displaying the three-dimensional graphic display, there may be generally used a program that is made so that a means for inputting the three-dimensional structural coordinate of the molecule, a means for measuring visual representation of the coordinate on a computer screen, the distance between the represented atoms in the molecule, bond angles or the like, a means for addition or modification of the coordinate, and the like can be provided. Furthermore, there may be used a program that has been made so that a means for calculating the structure energy of the molecule based on the

coordinate of the molecule, a means for calculating the free energy of bonding, and the ratio of bonding state number to non-bonding state number in consideration of solvent molecules such as water molecule can be provided. Examples of the program suitable for such purposes include Insight II, QUANTA and the like, which are computer programs commercially available from Accelrys Inc., and the present invention is not limited to these programs. Also, the above-mentioned programs can be introduced into a computer referred to as a work station supplied from Silicone Graphics Inc., SunMicro-Systems Ltd., or the like, and used.

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According to the crystal of dipeptidyl peptidase IV of the present invention, there can be exhibited excellent effects that the three-dimensional structural coordinate can be obtained as an information for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and that the crystal of a complex of the dipeptidyl peptidase IV and a known effector can be readily prepared. Also, according to the three-dimensional structural coordinate of the present invention, there is exhibited an excellent effect that the effector can be designed, identified, evaluated or searched. In addition, according to the method for obtaining a three-dimensional structural coordinate of the homolog protein of the dipeptidyl peptidase IV of the present invention, there is exhibited an excellent effect that the three-dimensional structural coordinate of the homolog protein of the dipeptidyl peptidase IV of which three-dimensional structure is unknown can be conveniently and rapidly provided. Furthermore, according to the method for

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obtaining a three-dimensional structure of a crystal of a complex of the dipeptidyl peptidase IV of the present invention and an effector of the dipeptidyl peptidase IV, there is exhibited an excellent effect that the method can provide a target for designing an effector useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and having excellent characteristics such as higher avidity, higher biological activity, higher biological stability, higher thermomechanical stability, and higher absorbency to a living body. Moreover, according to the method of the present invention for identifying a pharmacophore of the dipeptidyl peptidase IV and the effector of the dipeptidyl peptidase IV, there is exhibited an excellent effect that the method can provide a target for designing the effector useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and having excellent characteristics such as higher avidity, higher biological activity, higher biological stability, higher thermomechanical stability, and higher absorbency to a living body. According to the method of the present invention for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV, there is exhibited an excellent effect that the method can logically and conveniently provide an effector useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and having excellent characteristics such as higher avidity, higher biological activity, higher biological stability, higher thermomechanical stability,

and higher absorbency to a living body. According to the effector of the dipeptidyl peptidase IV of the present invention, there are exhibited excellent effects that the effector is capable of modifying immune response and capable of treating or preventing diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like. Furthermore, according to the program and medium therefor of the present invention, there is exhibited an excellent effect that the method for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV can be performed more rapidly and conveniently.

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The present invention will be hereinafter more specifically explained by the following Examples, but the present invention is not intended to be limited by the Examples in any way. Unless otherwise indicated, the reaction conditions, procedures and the like can be referred to the instruction manual attached to the reagents used, *Molecular Cloning A Laboratory Manual*, third edition, Sambrook et al. [issued by Cold Spring Harbor Laboratory Press (2001)], and the like.

Example 1 Construction of Recombinant Baculovirus for Expression of Soluble Human Dipeptidyl Peptidase IV

20 (1) Cloning of Soluble Human Dipeptidyl Peptidase IV (shDPPIV) cDNA

Caco-2 cells [provided by American Type Culture Collection (ATCC)]

were cultured at 37°C using Dulbecco's Modified Eagle Medium (manufactured by Invitrogen) containing 20% by volume of inactivated fetal bovine serum (manufactured by Invitrogen; inactivated by incubation at 56°C for 30 minutes)

and 1% by volume of nonessential amino acid (manufactured by Invitrogen), in

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the presence of 5% by volume of CO₂.

Next, total RNA was extracted from the Caco-2 cells obtained.

Extraction of the total RNA was carried out using a product manufactured by Nippon Gene Co. Ltd. under the trade name of ISOGEN in accordance with the attached instruction manual. The obtained total RNA was used as a template for RT-nested PCR method described below.

In order to obtain a nucleic acid corresponding to a soluble human DPPIV from which the signal peptide sequence was removed (amino acid nos: 33-766 of SWISS-PROT Accession No: P27487), first, a cDNA fragment sequence of human DPPIV gene was amplified by RT-nested PCR method with total RNA of the Caco-2 as a template.

The thermal profile in the PCR is 30 cycles of reaction, in which one cycle comprises denaturation at 94°C for 30 seconds, annealing at 55°C for 30 seconds and polymerase extension reaction at 72°C for 1 minute.

The amplified DNA fragment was separated by agarose gel electrophoresis method, and a small fragment of the gel of the corresponding band portions was cut out. Thereafter, the DNA fragment was extracted from the obtained small fragments of the gel using a product manufactured by Bio 101 under the trade name of GENE CLEAN SPIN Kit, and purified. The obtained fragment was inserted into vector pCR2.1-TOPO contained in TOPO TA Cloning (registered trade mark) Kit manufactured by Invitrogen to construct pCR-shDPPIV.

In order to confirm whether or not the obtained cDNA fragment encodes the desired polypeptide, deletion mutants regarding the DNA fragment having various lengths were prepared, and a nucleotide sequence for the DNA fragment

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was determined as follows.

First, a DNA fragment having a size of 2.2 kbp obtained by double digestion of the pCR-shDPPIV with BamHI and EcoRI was inserted into a corresponding restriction site in pUC19 (manufactured by Takara Bio Inc.), to construct a plasmid pUshDPPIV. Various deletion plasmids were prepared using the plasmid pUshDPPIV by a conventional method.

The nucleotide sequence for the DNA fragment was determined using the obtained deletion plasmid or plasmid pCR-shDPPIV, and a product manufactured by Perkin-Elmer Cetus Inc. under the trade name of Taq DyeDeoxy Terminator Cycle Sequencing Kit and Model 373S sequencer manufactured by Applied Biosystems.

Also, the amino acid sequence of the polypeptide encoded by the abovementioned DNA fragment was determined on the basis of the nucleotide sequence.

The determined amino acid sequence was compared with the sequence for a full length DPPIV of human colon shown in SEQ ID NO: 2. As a result, it was confirmed that the corresponding regions (regions excluding the transmembrane region) were identical.

Thus, it was confirmed that the DNA fragment encodes the desired polypeptide shDPPIV, namely a polypeptide in which the transmembrane region (amino acid nos: 1-32 at N-terminal side) in the full-length human DPPIV was deleted and a polyhistidine peptide was added to the C-terminal side.

(2) Preparation of Recombinant Baculovirus

Plasmid pUshDPPIV was digested with a restriction enzyme to give a

DNA fragment encoding shDPPIV gene. The obtained fragment was inserted into pAcGP67B (manufactured by BD PharMingen) to construct a baculovirus transfer vector pAcGP67B-shDPPIV.

Fifteen minutes before the transfection, Sf21 cells were washed twice with a TNM-FH medium comprising 10% by volume of fetal bovine serum. The Sf21 cells were then transferred to a well of a 6-well plate by 2.4×10^6 cells per well.

Furthermore, 2 to 5 µg of the baculovirus transfer vector and a 0.5 µg linear baculovirus DNA (trade name: BaculoGold virus DNA, manufactured by BD PharMingen) were mixed, and the mixture was allowed to stand at room temperature for 5 minutes. Next, 1 ml of Transfection Buffer B (manufactured by BD PharMingen) was added to the obtained mixture, and the mixture was thoroughly mixed to give a Transfection Buffer B/DNA mixture.

The medium in the wells of the 6-well plate and the cells that had not been adhered to the wells were removed, and 1 ml of Transfection Buffer A (manufactured by BD PharMingen) was added to each of the wells. The Transfection Buffer B/DNA mixture was gradually added dropwise to the wells of the 6-well plate, with gently stirring the 6-well plate. The cells were incubated at 28°C for 4 hours in the wells of the 6-well plate. Thereafter, the transfection buffer was removed, and 3 ml of TNM-FH medium containing 10% by volume of fetal bovine serum was added to the wells of the 6-well plate. The cells were cultured at 28°C in each of the wells of the 6-well plate for 5 days, and the culture supernatant was collected. The culture supernatant was used for amplification of virus using Sf21 cells to give a virus stock solution.

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Example 2 Preparation and Crystallization of shDPPIV

(1) Expression of shDPPIV in Insect Cells

(manufactured by JRH Biosciences) and T flask, and Tn5 cells (provided by Invitrogen) were cultured using a serum free medium EX-CELL 401 (manufactured by JRH Biosciences) and a T flask, at 28°C, respectively. At the time when the proliferation of the cells reached 70% confluent, the old medium was removed, and a fresh medium was added at 40 ml per one 225-cm² flask. Then, 1.5 ml of virus stock solution after amplification for three times (having multiplicity of infection (MOI) of about 2) was added to the cells to infect the cells, and the cells were incubated at 28°C for 4 days. The culture supernatant four days after the infection was collected and stored at -20°C. The culture supernatant was used for the purification of shDPPIV protein as described below.

15 (2) Purification of shDPPIV Protein

In each step for the purification of shDPPIV, the activity of DPPIV was measured by incubating a 0.1 ml reaction mixture containing a 1.5 mM substrate [manufactured by Peptide Institute, Gly-Pro-paranitroanilide (pNA)], 71 mM Gly-NaOH (pH 8.7) and the DPPIV, and detecting the liberated pNA.

Meanwhile, the reaction mixture was incubated at 37°C for 10 minutes. During the incubation, the absorbance at 405 nm was monitored.

Also, the protein concentration was quantified by using a product manufactured by Bio-Rad Laboratories, Inc. under the trade name of DC protein Assay Kit II.

The purity of the protein was confirmed by subjecting a protein sample

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in each step to SDS-PAGE using 7.5% polyacrylamide gel according to the method by Laemmli et al.

The culture supernatant stored at -20°C in the above-mentioned (1) was melted at 4°C and filtered with a bottle top filter (manufactured by Becton, Dickinson and Company) or with 0.45 µm filter (KURABO INDUSTRIES LTD.) to remove insoluble materials. The supernatant after the filtration was concentrated to an about tenth volume by using a concentrator Vivaflow 50 (manufactured by Sartorius AG) or Amicon stirrer cell model 8400 (manufactured by Millipore Corporation) to give a concentrated solution.

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The obtained concentrated solution was dialyzed against buffer A (20 mM HEPES-NaOH, 0.5 M NaCl, pH 8.0) at 4°C overnight, and applied to a nickel column [one in which nickel was immobilized to HiTrap Chelating column (trade name, manufactured by Amersham-Pharmacia) (5 ml × 2)] equilibrated with buffer A. The column was washed with 10 column volumes of buffer A, and then with buffer A containing 50 mM imidazole. The elution of the fraction containing shDPPIV was carried out by a linear gradient of 50 to 500 mM imidazole. The fraction found to have DPPIV activity was collected, and dialyzed overnight at 4°C against buffer B (20 mM HEPES-NaOH, pH 8.0, 50 mM NaCl). After the dialysis, the sample was purified by using an anion exchange column [manufactured by Amersham-Pharmacia under the trade name: Resource Q (6 ml)] equilibrated with buffer B. The column was washed with buffer B, and thereafter shDPPIV was eluted by a linear gradient of 15 column volumes of 50 to 500 mM NaCl. The fractions found to have DPPIV activity were collected, and used as a purified preparation.

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(3) Preparation of Protein Sample for Crystallization

The shDPPIV purification sample (9 ml) obtained in the above (2) was concentrated using a product manufactured by Millipore Corporation under the trade name of Centricon 10 until the protein concentration reached 10 mg/ml.

The obtained product was used as a protein sample for crystallization. The protein sample for crystallization was stored at 4°C.

A precipitation agent solution containing 0.18 M glycine-NaOH (pH 9.5), 0.18 M sodium sulfate and 18% by weight of PEG4000, and a 10 mg/ml dipeptidyl peptidase IV solution were mixed, and thereafter, a drop of the obtained mixed solution was placed on a product under the trade name of Cryschem Plate (manufactured by Hampton Research). The above-mentioned precipitation solution was allowed to stand at 20°C as a reservoir solution to allow crystallization.

15 (4) Crystallization of shDPPIV

The crystallization of shDPPIV was carried out by a sitting-drop method, which is one of vapor diffusion methods.

The formation of crystal was observed with the passage of time using a stereoscopic microscope. As a result, after about two weeks, a large crystal having a maximum size of 500 $\mu m \times 300~\mu m \times 100~\mu m$ was obtained. The crystal is also referred to as a native crystal. The microphotograph of the obtained crystal is shown in Figure 1. In Figure 1, the visual field is $4000~\mu m \times 3000~\mu m$.

25 Example 3 Three-Dimensional Structural Analysis of Crystals

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(1) X-ray Diffraction

The crystal obtained in Example 2 mentioned above was soaked in a cryoprotecting buffer [composition: 0.18 M glycine-NaOH (pH 9.5), 19% by weight of PEG4000, 0.18 M sodium acetate, 15% glycerol], and immediately thereafter the mixture was placed under nitrogen gas stream (100 K) to rapidly freeze the mixture.

The X-ray diffraction intensity data of the above crystal were collected up to the resolution of 3.0Å using a product manufactured by Rigaku International Corporation under the trade name of R-AXIS IV in nitrogen gas stream (100 K), and converted to the structure factor using a program MOSFLM (Version 6.11). A photograph of the diffraction pattern is shown in Figure 2.

From the obtained diffraction intensity data, it was determined that the crystal form to which the crystal belongs was orthorhombic, that the space group was $P2_12_12_1$, and the lattice constants were $a = 118.0 \pm 5.0$ Å, $|b| = 125.9 \pm 5.0$ Å and $|c| = 136.8 \pm 5.0$ Å.

(2) Multiple Isomorphous Replacement Method

In order to derive an electron density map, multiple isomorphous replacement method was carried out. The crystal obtained in Example 2 was soaked for 3 days and 4 days in a crystallization solution prepared by dissolving mercury chloride until being saturated, to give two different kinds of isomorphous replacement crystals containing mercury atoms in the crystals. The X-ray diffraction intensity data were collected in the same manner as those for the native crystals.

In the determination of the phase in the structural analysis, CCP4

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(Collaborative Computational Project, Number 4, 1994. "The CCP4 Suite: Programs for Protein Crystallography," *Acta Cryst.* D50, 760-763) program was used.

First, Fourier transform calculation utilizing the difference between the diffraction intensity obtained from the two kinds of isomorphous replacement crystals of mercury and the diffraction intensity obtained from the native crystal was performed using MLPHERE contained in the CCP program package. The position of each mercury atom in the unit cell of the real space was determined by investigating large peaks provided by heavy atoms (mercury) in the obtained Patterson's diagram. The phase of the crystal structure factor of the native crystals was determined by using the obtained position coordinate of mercury atoms. Furthermore, in order to determine the coordinate of each mercury atom more accurately using DM and SOLOMON contained in the CCP program package, refinement was carried out using three crystal structure factors of the native crystals and of the two kinds of mercury isomorphous replacement crystals.

An electron density map of the crystals of the dipeptidyl peptidase IV in real space was obtained using the phase of the crystal structure factor of the native crystals calculated from the refined coordinates of the mercury atoms. Furthermore, the electron density map was improved by carrying out smoothening and histogram matching of the electron density map in a solvent region, to obtain an electron density map critical for molecular modeling.

(3) Molecular Modeling

25 The sites corresponding to the amino acid residues of the dipeptidyl

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peptidase IV were identified on the electron density map by using QUANTA (manufactured by Accelrys, Inc.), to build molecular models.

As expected from the lattice constants, there were two molecules of the dipeptidyl peptidase IV in an asymmetry unit, and a model was built for each of the molecules. The refinement of the obtained molecular model was carried out using CNX (manufactured by Accelrys, Inc.), and the molecular model was adjusted again using the QUANTA for the obtained improved electron density map. The procedures were repeated to build a more accurate molecular model. In the refinement of the final coordinate, diffraction intensity data measured again were used after OSMIC confocal mirror (manufactured by Rigaku International Corporation) had been introduced into R-AXIS IV (trade name, manufactured by Rigaku International Corporation).

As a result, the resolution was improved from the previous 3.0Å to 2.6Å. Furthermore, 273 molecules of bound water and 5 molecules of N-acetyl glucosamine residues per molecule of the dipeptidyl peptidase IV were identified in an asymmetric unit. R factor, which is an index for accuracy of the obtained molecular model, was 24.89%, and a free R factor, which independently was not taken into account of the calculation of refinement at the step of refinement, was 30.15%. During the procedure, the deviation of the interatomic bond distance (rms-deviation) and the bond angle from the ideal state of the three-dimensional structure were 0.006Å and 1.305°, respectively. The stereogram of the three-dimensional structure model of the crystals is shown in Figure 3, and the coordinate is shown in Figure 4. The present coordinate data were registered in PDB (Brookhaven Protein Data Bank) [PDB Code No: 1J2E, RSCB code No: 005544].

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Here, as to those regions which did not take a regular structure in the crystals (in the disordered state), namely, the region from Asp 38 to that closer to the N-terminal side thereof, and the region for the tagged peptide (polyhistidine peptide) of the C-terminal side, the molecular model could not be built.

Furthermore, a part of the side chains projected to the surface of the molecules did not take a regular structure. However, these residues were not portions that play an important role for the function of enzymes.

In the three-dimensional structure of the dipeptidyl peptidase IV, which has been clarified by the Examples, it has been revealed that the amino acid residue involved in the activity deduced by various experiments for the dipeptidyl peptidase IV, namely, Ser 630, Asp 708 and His 740, form hydrogen bonds between the $O_{\delta 2}$ atom of Asp 708 and $N_{\delta 1}$ atom of His 740, and with the $N_{\epsilon 2}$ atom of His 740 and O_{γ} atom of Ser 630, even the residues locate in distant locations on the primary sequence. Therefore, for the structural coordinate of Figure 4 and the three-dimensional structure model defined by the structural coordinate, it is suggested that the regions characterized by Ser 630, Asp 708 and His 740, and the whole or a part of amino acid residues that are located in the vicinity of Ser 630, Asp 708 and His 740 play an important role on the exhibition of the activity for the dipeptidyl peptidase IV and binding or interaction of the dipeptidyl peptidase IV with the effector, and that the compound matching the three-dimensional structure of the regions affect the activity for the dipeptidyl peptidase IV.

The present invention may be embodied in other various forms without departing from the spirit or essential characteristics thereof. The present

embodiment is therefore to be considered in all respects as illustrative and not restrictive, the scope of the invention being indicated by the appended claims rather than by the foregoing description and all changes which come within the meaning and range of equivalency of the claims are therefore intended to be embraced therein.

INDUSTRIAL APPLICABILITY

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According to the crystal of the dipeptidyl peptidase IV of the present invention, the information of a three-dimensional structure coordinate suitable for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV, useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like can be obtained. Also, according to the three-dimensional structure coordinate, the information of a three-dimensional structure coordinate suitable for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV, useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like can be obtained. Further, according to the method of the present invention for obtaining a threedimensional structure coordinate of a homolog protein of a dipeptidyl peptidase IV, the refinement of the three-dimensional structure coordinate of the homolog protein of the dipeptidyl peptidase IV can be more conveniently carried out. Moreover, according to the method of the present invention for obtaining a threedimensional structure coordinate of a crystal of a complex of a dipeptidyl

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peptidase IV with an effector of the dipeptidyl peptidase IV, the information of a three-dimensional structure coordinate suitable for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV, which is useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and is excellent in avidity, biological activity, biological stability, absorbency to a living body, and which can favorably act on the dipeptidyl peptidase IV can be obtained. Also, according to the method for identifying a pharmacophore of a dipeptidyl peptidase IV and an effector of the dipeptidyl peptidase, the information of a three-dimensional structure coordinate suitable for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV, which is useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and is excellent in avidity, biological activity, biological stability, absorbency to a living body, and which can favorably act on the dipeptidyl peptidase IV can be obtained. Further, according to the method of the present invention for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV, the information of a three-dimensional structure coordinate suitable for designing, identifying, evaluating or searching an effector of the dipeptidyl peptidase IV, which is useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like, and is excellent in avidity, biological activity, biological stability, absorbency to a living body,

and which can favorably act on the dipeptidyl peptidase IV can be logically and conveniently obtained. In addition, the effector of the dipeptidyl peptidase IV of the present invention is useful as a modulatory agent of immune response and as a therapeutic or prophylactic agent for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like. Further, according to the program or the medium therefor of the present invention, the design, identification, evaluation and search for an effector of a dipeptidyl peptidase IV can be carried out rapidly and conveniently. Therefore, the present invention can be utilized in modulation of immune response and the treatment or prevention for diabetes, inflammation, multiple sclerosis, Graves' disease, chronic rheumatoid arthritis, AIDS, cancer and the like.

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CLAIMS

- 1. A crystal of a dipeptidyl peptidase IV, having characteristics sufficient to ensure a resolution capable of analyzing its three-dimensional structure up to the side chain level by X-ray crystallographic structural analysis.
- 2. The crystal according to claim 1, wherein the dipeptidyl peptidase IV is a soluble polypeptide comprising a region located at extramembrane in a full-length human dipeptidyl peptidase IV.

3. The crystal according to claim 1 or 2, wherein the dipeptidyl peptidase IV is a polypeptide having an amino acid sequence in which a transmembrane region is deleted from the amino acid sequence of SEQ ID NO: 2, and a tag peptide is optionally added to a C-terminal side or N-terminal side thereof.

4. The crystal according to any one of claims 1 to 3, wherein the crystal has a space group of $P2_12_12_1$, and a lattice constant of the unit cell of $|a| = 118.0 \pm 5.0 \text{Å}$, $|b| = 125.9 \pm 5.0 \text{Å}$, $|c| = 136.8 \pm 5.0 \text{Å}$, and $\alpha = \beta = \gamma = 90^{\circ}$, and is orthorhombic.

- 5. The crystal according to any one of claims 1 to 4, wherein the crystal has the structural coordinate shown in Figure 4.
- 6. The crystal according to any one of claims 1 to 4, wherein the crystal has a structural coordinate different from the structural coordinate as shown in

Figure 4 via fluctuation of a protein.

7. A three-dimensional structural coordinate of a dipeptidyl peptidase IV, comprising the structural coordinate shown in Figure 4.

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- 8. A three-dimensional structural coordinate of a dipeptidyl peptidase IV, comprising a structural coordinate different from the structural coordinate as shown in Figure 4 via fluctuation of a protein.
- 9. The three-dimensional structural coordinate according to claim 8, wherein the fluctuation of a protein is a state that is caused by molecular oscillation or temperature, and exhibits an activity for the dipeptidyl peptidase IV in a living body.
- 15 10. The three-dimensional structural coordinate according to any one of claims 7 to 9, wherein the dipeptidyl peptidase IV is a soluble polypeptide comprising a region located at extramembrane in a full-length human dipeptidyl peptidase IV.
- 20 11. The three-dimensional structural coordinate according to any one of claims 7 to 10, wherein the dipeptidyl peptidase IV is a polypeptide having an amino acid sequence in which a transmembrane region is deleted from the amino acid sequence of SEQ ID NO: 2, and a tag peptide is optionally added of to a C-terminal side or N-terminal side thereof.

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- 12. A three-dimensional structural coordinate of a region in a dipeptidyl peptidase IV, comprising the three-dimensional structural coordinate of the region selected from the group consisting of the following (a) to (d):
- (a) a region characterized by Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and all or a part of a group of the amino acid residues located in the adjacent area of each of the Ser 630, Asp 708 and His 740 in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate;
- 10 (b) a region characterized by Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and all or a part of a group of the amino acid residues comprising amino acids capable of maintaining physicochemical characteristics physiologically equivalent to each of amino acids in the group of the amino acid residues located in the adjacent area of each of Ser 630, Asp 708 and His 740, in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate,
 - (c) a region characterized by a group of amino acid residues comprising amino acids capable of maintaining physicochemical characteristics physiologically equivalent to each of Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and all or a part of a group of the amino acid residues located adjacent area of said group of the amino acid residues in the structural coordinate shown in Figure 4 or the three-dimensional structure model defined by the structural coordinate; and

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- (d) a region characterized by a group of amino acid residues comprising amino acids capable of maintaining physicochemical characteristics physiologically equivalent to each of Ser 630, Asp 708 and His 740 in the amino acid sequence of SEQ ID NO: 2, and all or a part of a group of amino acid residues comprising amino acids capable of maintaining physicochemical characteristics physiologically equivalent to each of the amino acids in the group of the amino acid residues located in the adjacent area of said group of the amino acids, in the structural coordinate shown in Figure 4 or the three-dimensional
- wherein the region in the dipeptidyl peptidase IV is a region involved in binding or interaction between the dipeptidyl peptidase IV and an effector of the dipeptidyl peptidase IV.

structure model defined by the structural coordinate,

- 13. The three-dimensional coordinate according to claim 12, wherein the physicochemical characteristic is selected from the group consisting of features in shape of a three-dimensional structure, hydrophobicity, electric charge and pK.
- 14. A method for obtaining a three-dimensional coordinate of a homolog

 protein of a dipeptidyl peptidase IV, characterized in refining an electron density
 map of the homolog protein of the dipeptidyl peptidase IV comprising the amino
 acid sequence of SEQ ID NO: 2, based on all and/or a part of the threedimensional coordinate of any one of claims 7 to 13, to give a three-dimensional
 structural coordinate.

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- 15. A method for obtaining a three-dimensional structural coordinate of a crystal of a complex of a dipeptidyl peptidase IV and an effector of the dipeptidyl peptidase IV characterized in using all and/or a part of the three-dimensional structural coordinate of any one of claims 7 to 13, to give a three-dimensional structural coordinate.
- 16. A method for identifying pharmacophore of an effector of the dipeptidyl peptidase IV, characterized in identifying the pharmacophore based on all and/or a part of the three-dimensional structural coordinate of any one of claims 7 to 13, and the steric conformation of the effector of the dipeptidyl peptidase IV.
- 17. A method for designing, identifying, evaluating or searching an effector of a dipeptidyl peptidase IV, characterized in designing, identifying, evaluating or searching a compound capable of acting on the dipeptidyl peptidase IV, based on all and/or a part of the three-dimensional structural coordinate of any one of claims 7 to 13.
- 18. The method according to claim 17, wherein the method for designing, identifying, evaluating or searching an effector comprises the steps of:
- 20 (i) identifying a region to be targeted for binding or interaction with the effector in a dipeptidyl peptidase IV, based on all and/or a part of the three-dimensional structural coordinate according to any one of claims 7 to 13 and the steric conformation of the effector of the dipeptidyl peptidase IV;
- 25 (ii) identifying atoms or atomic groups capable of generating in the above

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region at least one intermolecular interaction selected from the group consisting of covalent bond, ionic interaction, ion-dipole interaction, dipole-dipole interaction, hydrogen bonding, van der Waals force, electrostatic interaction and hydrophobic interaction, with the atoms or atomic groups existing in a candidate compound; and

- (iii) designing a compound based on the information of the above step (i) and/or (ii).
- 19. The method according to claim 18, wherein the method further comprisesthe steps of:

detecting an interaction between the dipeptidyl peptidase IV and the designed, identified, evaluated or searched candidate compound, wherein when an interaction is detected, the candidate compound is identified as a compound capable of binding to the dipeptidyl peptidase IV, based on a degree of the interaction as an index.

20. The method according to claim 18 or 19, wherein the method further comprises the steps of:

contacting the dipeptidyl peptidase IV with the designed, identified,

evaluated or searched candidate compound and measuring an activity of the
dipeptidyl peptidase IV,
wherein when an activity increases or decreases, the designed, identified,
evaluated or searched candidate compound is identified as a compound having
enhancing action or inhibitory action on the activity of the dipeptidyl peptidase

IV, based on a degree of the increase or decrease as an index.

- 21. An effector of the dipeptidyl peptidase IV obtainable by the method of any one of claims 17 to 20.
- 5 22. A program and a medium therefor for use of the three-dimensional structural coordinate of any one of claims 7 to 13, wherein all and/or a part of the three-dimensional structural coordinate of any one of claims 7 to 13 is recorded.
- 23. The program and the medium according to claim 22, comprising a means for identifying, searching, evaluating or designing a compound capable of binding to the dipeptidyl peptidase IV or a compound having an enhancing action or inhibitory action on the activity for the dipeptidyl peptidase IV.
- 24. The program and the medium according to claim 23, further comprising a means for displaying a three-dimensional graphic display of a molecule.

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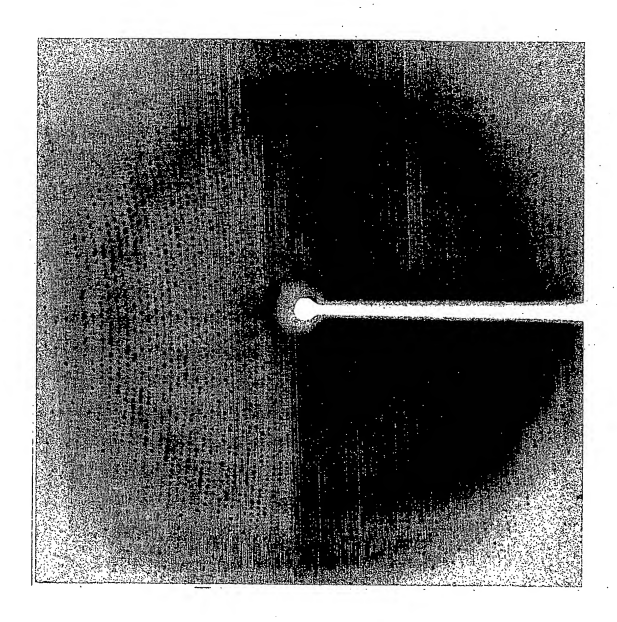
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FIG. 1



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FIG. 2



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FIG. 3

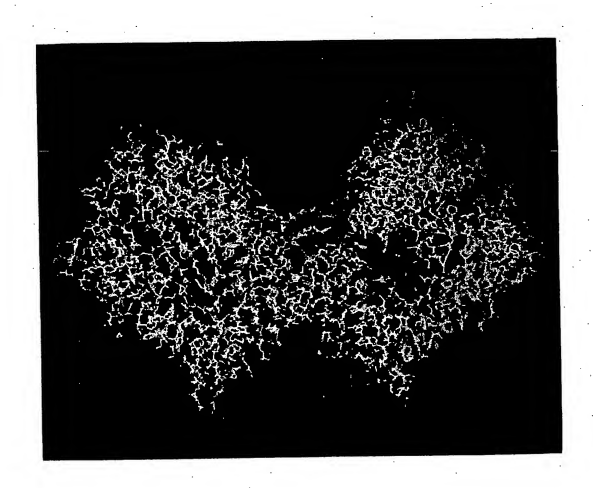


FIG. 4-1

]	Chree	-dime	nsional	structural	coordi	nate of	dipept	idyl	peptidase	IV
ATOM											
ATOM	$\frac{1}{2}$	CB CG	ASP	38	44. 493	31.885			42.4		C ·
ATOM	3		ASP	38	44. 146	32. 095			42.0		C
ATOM			ASP ASP	38	43.664	33. 198			42.5		0
ATOM	4			38	44. 360	31.171			40.8		0
ATOM	5	C	ASP	38	45. 876	29.805			41.6		C
ATOM	. 6 7	O N	ASP	38	46. 980	30. 327			42.0		0
ATOM	8		ASP	38	44. 758	30. 264			42.8		N
ATOM	9	CA N	ASP SER	38	44. 639	30.404	59. 296		42.5		C
ATOM	10	CA	SER	39 39	45.679	28. 711			40.6		N
ATOM	11	CB	SER	39 39	46.775	28. 013	57. 241		39.9	_	C
ATOM	12	OG	SER	39 39	46. 584	26. 501	57. 380		40.4		C
ATOM	13	C	SER	39 39	45.410	26.079	56. 703		41.1		0
ATOM	14	0	SER	39	46.960	28. 343	55. 763		39.6		C
ATOM	15	N	ARG	39 40	47.870	27. 813	55. 123		39.6		0
ATOM	16	CA	ARG	40	46. 093 46. 194	29. 190	55. 217		38.1		N
ATOM	17	CB	ARG	40	45. 082	29.575	53.810		37.0		C
ATOM	18	CG	ARG	40	43. 683	30. 558 29. 984	53. 439		36.9	-	C
ATOM	19	CD	ARG	40	42. 688	31.098	53. 404 53. 137		35.9		C
ATOM	20	NE	ARG	40	42. 774	32. 134	54. 161		34.9	_	C
ATOM	21	CZ	ARG	40	42.097	33. 27.6	54. 125		35. 2		N
ATOM	22	NH1	ARG	40	41. 280	33. 528	53. 111		35. 54 35. 54		C
ATOM	23		ARG	40	42. 239	34. 167	55. 097		34. 68		N
ATOM	24	C	ARG	40	47. 530	30. 251	53. 531		35. 91		N
ATOM	25	0	ARG	40	48. 100	30. 901	54. 407		34. 18	_	C
ATOM	26	N	LYS	41	48. 031	30. 100	52. 310		35. 43		0
ATOM	27	CA	LYS	41	49. 286	30. 749	51. 937		34. 97		N
ATOM	28	CB	LYS	41	49. 705	30. 338	50. 525		35. 73		C
ATOM	29	CG	LYS	41	48. 684	30.719	49. 467		38. 56		Č
ATOM	30	CD	LYS	41	49.026	30. 151	48. 096		42. 36		Č
ATOM	31	CE	LYS	41	47.805	30. 201	47. 173		45. 55		Č
ATOM	32	NZ	LYS	41	48.070	29.686	45. 791		47.41		N
ATOM	33	C	LYS	41	49.038	32.257	51.957		33. 41		Č
ATOM	34		LYS	41	47.891	32.715	51.981		33. 24		ŏ
ATOM	35	N	THR	42	50.110	33.032	51.954		31.47		N
ATOM	36	CA	THR	42	49.967	34.479	51.937				Č
ATOM	37		THR	42	50.860	35.139	53.000		31. 23	Ä	č
ATOM	38		THR	42	52. 234	34.843	52.725		30.79		ŏ
ATOM	39	CG2		42	50. 501	34.622	54.386		30.12		Č
ATOM	40		THR	42	50. 389	34.971	50.558		28.34		Č
ATOM	41			42	50. 977	34. 220	49.782		27.76		ŏ
ATOM	42		TYR	43	50.058	36.217	50. 234		27.55		Ň
ATOM	43		TYR	43	50.465	36. 782	48.954		25.72		Ċ
ATOM	44		TYR	43	49.615	38.006	48.623	1.00		Ä	Č
ATOM	45		TYR	43		38.625	47.280	1.00		A	č
ATOM	46		TYR	43		39.527	47.130	1.00		A	Č
ATOM	47		TYR	43		40.113	45.895	1.00		Ā	Č
ATOM	48	CD2	IYK	43	49. 152	38. 315	46.158	1.00		A	Č

				(Continued)
•			FIG. 4-2	
ATOM	49 CE2 TYR	43	49. 424 38. 891 44. 919 1. 00 25. 89	A C A C
ATOM	50 CZ TYR	43	50. 473 39. 790 44. 796 1. 00 25. 91 50. 741 40. 370 43. 579 1. 00 25. 09	A O
ATOM	51 OH TYR	43		A C
ATOM	52 C TYR 53 O TYR	43 43	51. 933 37. 165 49. 160 1. 00 24. 97 52. 251 38. 049 49. 955 1. 00 23. 33	A O
ATOM	53 0 TYR 54 N THR	43 44	52. 818 36. 482 48. 444 1. 00 24. 06	A N
ATOM ATOM	55 CA THR	44	54. 255 36. 685 48. 580 1. 00 25. 90	A C
ATOM	56 CB THR	44	54.960 35.336 48.547 1.00 25.86	A C
ATOM	57 OG1 THR	44	54, 696 34, 709 47, 285 1, 00 28, 12	A 0
ATOM	58 CG2 THR	44	54. 439 34. 436 49. 655 1. 00 22. 61	A C
ATOM	59 C THR	44	54.917 37.576 47.530 1.00 27.35	A C
ATOM	60 0 THR	44	54. 296 37. 956 46. 535 1. 00 29. 11	A O
ATOM	61 N LEU	45	56. 191 37. 894 47. 765 1. 00 27. 39	A N
ATOM	62 CA LEU	45	56. 978 38. 722 46. 853 1. 00 26. 43	A C
ATOM	63 CB LEU	45	58. 377 38. 954 47. 425 1. 00 26. 07 59. 310 39. 860 46. 612 1. 00 26. 21	A C A C
ATOM	64 CG LEU	45	59. 310 39. 860 46. 612 1. 00 26. 21 58. 734 41. 263 46. 517 1. 00 25. 53	A C A C A C
ATOM	65 CD1 LEU 66 CD2 LEU	45 45	60.672 39.896 47.266 1.00 24.37	A Č
ATOM ATOM	67 C LEU	45	57. 088 38. 069 45. 473 1. 00 27. 00	A C
ATOM	68 0 LEU	45	56. 939 38. 740 44. 449 1. 00 27. 84	A 0
ATOM	69 N THR	46	57. 354 36. 766 45. 445 1. 00 26. 70	A N
ATOM	70 CA THR	46	57.448 36.038 44.182 1.00 26.95	A C
ATOM	71 CB THR	46	57. 838 34. 559 44. 407 1. 00 26. 87	A C
ATOM	72 OG1 THR	46	59. 150 34. 495 44. 966 1. 00 31. 74	A 0
ATOM	73 CG2 THR	. 46	57. 833 33. 793 43. 110 1. 00 28. 08	A C A C
ATOM	74 C THR	46	56.076 36.091 43.517 1.00 26.96 55.965 36.094 42.289 1.00 25.36	A C A O
ATOM	75 0 THR 76 N ASP	46 47	55.965 36.094 42.289 1.00 25.36 55.035 36.126 44.346 1.00 27.72	A N
ATOM ATOM	76 N ASP 77 CA ASP	47	53.659 36.199 43.858 1.00 29.74	A C
ATOM	78 CB ASP	47	52.670 36.173 45.026 1.00 30.90	A C
ATOM	79 CG ASP	47	52. 289 34. 769 45. 430 1. 00 30. 62	A C
ATOM	80 OD1 ASP		51.778 34.595 46.553 1.00 32.28	A 0
ATOM	81 OD2 ASP	47	52.490 33.845 44.617 1.00 30.71	A 0
ATOM	82 C ASP		53. 477 37. 482 43. 073 1. 00 28. 87	, A C
ATOM	83 0 ASP		52. 918 37. 478 41. 979 1. 00 29. 50	A O
ATOM	84 N TYR		53. 945 38. 581 43. 648 1. 00 28. 54	A N A C
ATOM	85 CA TYR		53. 859 39. 878 42. 994 1. 00 29. 04 54. 191 40. 991 43. 996 1. 00 27. 50	A C
ATOM	86 CB TYR 87 CG TYR		54.448 42.333 43.354 1.00 25.16	A Č
ATOM ATOM	87 CG TYR 88 CD1 TYR		53. 460 42. 971 42. 609 1. 00 23. 19	Ä Č .
ATOM	89 CE1 TYR		53. 703 44. 184 41. 982 1. 00 24. 84	A C
ATOM	90 CD2 TYR		55.694 42.946 43.461 1.00 25.89	A C
ATOM	91 CE2 TYR		55.956 44.165 42.838 1.00 26.76	A C
ATOM	92 CZ TYR		54. 955 44. 779 42. 096 1. 00 27. 28	A C
ATOM	93 OH TYR	48	55. 208 45. 977 41. 463 1. 00 25. 97	A 0
ATOM	94 C TYR		54. 820 39. 953 41. 796 1. 00 28. 80	A C
ATOM	95 0 TYR		54. 445 40. 401 40. 714 1. 00 28. 24	A O
ATOM	96 N LEU		56.054 39.499 41.988 1.00 29.41 57.046 39.552 40.918 1.00 30.39	A N A C
ATOM	97 CA LEU	J 49	57.046 39.552 40.918 1.00 30.39	n v

					F]	G. 4	: - 3			(Continued)
ATOM ATOM	98 99	CB CG	LEU LEU	49 49	58. 455 58. 988	39.318 40.473	41. 481 42. 336	1.00 27.73 1.00 28.28	A A	C C
ATOM	100	CD1		49	60.438	40. 223	42. 711	1.00 26.29	A	č
ATOM	101		LEU	49	58. 860	41.773	41.555	1.00 26.02	Ä	č
ATOM	102	C	LEU	49	56.804	38.606	39.752	1.00 30.71	A	č
ATOM	103	0	LEU	49	57.147	38.919	38.614	1.00 30.14	· Ā	0
ATOM	104	N	LYS	50	56.198	37.459	40.024	1.00 32.51	Ā	N
ATOM	105	CA	LYS	50	55.959	36.491	38.971	1.00 33.54	A	Č .
ATOM	106	CB	LYS	50	56.,289	35.098	39.485	1.00 33.30	Α	С
ATOM	107	CG	LYS	50	57.763	34.940	39.790	1.00 33.89	Α	C
ATOM	108	CD	LYS	50	58.591	35. 213	38. 545	1.00 35.19	Α	С
ATOM	109	CE	LYS	50	60.071	34. 945	38. 778	1.00 38.12	Α	C
ATOM	110	NZ	LYS	50	60.859	35.028	37. 515	1.00 39.27	Α	N
ATOM	111	C	LYS	50	54. 572	36.517	38. 361	1.00 34.93	Α	· C
ATOM	112	0	LYS	50	54. 272	35.719	37. 478	1.00 35.13	Α	0
ATOM	113	N ·	ASN	51	53. 731	37. 436	38.822	1.00 36.66	A	N
ATOM	114	CA	ASN	51	52. 379	37.569	38. 294	1.00 38.39	A	C
ATOM	115	CB	ASN	51	52. 428	37.859	36. 791	1.00 41.61	A	C
ATOM	116	CG	ASN	51	53. 407	38.968	36. 436	1.00 44.75	A	C
ATOM	117 118		ASN	51	53. 212	40.131	36.801	1.00 46.38	A	0
ATOM ATOM	119	C	ASN ASN	51 51	54. 470	38.609	35.717	1.00 45.80	A	N
ATOM	120	0	ASN	51 51	51. 529 50. 708	36. 324 35. 976	38. 517 37. 674	1.00 38.21 1.00 40.60	A	C
ATOM	121	N	THR	52	51.720	35.647	39. 641	1.00 40.00	A	0 N
ATOM	122	CA	THR	52 52	50. 942	34. 451	39. 926	1.00 35.44	A A	C ·
ATOM	123	CB	THR	52 52	51. 297	33. 888	41. 298	1.00 35.44	A	Č
ATOM	124	0G1	THR	52	52.646	33.415	41. 272	1.00 38.62	A	Õ
ATOM	125	CG2		52	50. 367	32.750	41.666	1.00 35.25	A	
ATOM	126	C	THR	52	49. 431	34.686	39. 869	1.00 35.17	A	C
ATOM	127	Ŏ	THR	52	48.699	33. 889	39. 276	1.00 36.44	. A	ŏ
ATOM	128	N	TYR	53	48.962	35. 765	40. 487	1.00 33.55	Ä	Ň
ATOM	129	CA	TYR	53	47. 535	36.081	40.487	1.00 33.46	A	Ċ
ATOM	130	CB	TYŘ	53	47.084	36.407	41.903	1.00 32.64	Ā	Č
ATOM	131	CG	TYR	53	47.399	35. 293	42.861	1.00 33.83	A	Č
ATOM	132	CD1	TYR	53	48.341	35.462	43.872	1.00 34.11	Α	C
ATOM	133	CE1	TYR	53	48.657	34.425	44. 741	1.00 34.24	Α	C
ATOM	134	CD2		53	46.775	34.050	42.741	1.00 36.17	Α	C .
ATOM	135	CE2	TYR	53	47.084	33.001	43.605	1.00 35.64	Α	C
ATOM	136	CZ	TYR	53	48.026	33. 199	44.601	1.00 35.74	Α	C
ATOM	137	OH	TYR	53	48. 343	32.170	45.453	1.00 35.79	A	0 .
ATOM	138	C	TYR	53	47.266	37. 248	39. 548	1.00 33.40	Α	С
ATOM	139	0	TYR	53	47. 486	38.404	39.895	1.00 33.56	Α	0.
ATOM	140	N	ARG	54	46.773	36. 929	38. 355	1.00 34.36	A	Ŋ.
ATOM	141	CA	ARG	54	46.526	37. 933	37. 327	1.00 34.87	A	C
ATOM	142	CB	ARG	54	46. 993	37. 387	35.972	1.00 35.72	. A	Ċ
ATOM	143	CG	ARG	54	46.887	38. 373	34. 821	1.00 39.96	A	C
ATOM	144	CD	ARG	54	47.675	37. 880	33. 613	1.00 43.22	A	. <u>C</u>
ATOM	145	NE CZ	ARG	54 54	47.651	38.831	32. 506	1.00 46.70	A	N
ATOM	146	CZ	ARG	54	46. 587	39.068	31.744	1.00 49.10	A	С

				,						(Continued)
	•			_	· FI	G. 4	- 4			(Continued)
				•	T. I	U. I	•			
ATOM	147	NH1	ARG	54	45.451	38.416	31.968	1.00 49.25	A	N
ATOM	148	NH2	ARG	54	46.657	39.957	30. 757	1.00 50.00	A	N
ATOM	149		ARG	54	45.100	38. 445	37. 202	1.00 33.84	A	C
ATOM	150		ARG	54	44. 141	37. 687	37. 314	1.00 34.59	A	0 N
ATOM	151		LEU	55	44.982	39.748	36. 966	1.00 33.05	A	N C
ATOM	152		LEU	55	43.693	40. 402	36. 788	1.00 32.40 1.00 29.74	A A	C
ATOM	153		LEU	55	43. 792	41.892	37.123	1.00 29.74	A	C
ATOM	154		LEU	55	44.042	42.344	38. 557 38. 571	1.00 32.20	A	Č
ATOM	155	CD1		55	44. 245	43.847	39. 448	1.00 33.66	A	č
ATOM	156	CD2		55 55	42. 857 43. 298	41.967 40.271	35. 322	1.00 33.60	A	Č .
ATOM	157	C	LEU	55 55	43. 296	40. 769	34. 441	1.00 33.62	Ä	Ŏ
ATOM	158	0 N	LEU LYS	55 56	42.189	39. 593	35.050	1.00 31.32	Ä	Ň
ATOM	159	N CA	LYS	56	41. 733	39.462	33.673	1.00 31.42	A	Ċ
ATOM ATOM	160 161	CB	LYS	56	40.584	38. 453	33. 564	1.00 33.54	A	
ATOM	162	CG	LYS	56	40.978	36. 997	33. 733	1.00 34.84	Α	С
ATOM	163	CD	LYS	56	41.746	36. 484	32. 530	1.00 38.85	Α	C C C C
ATOM	164	CE	LYS	56	42.120	35.009	32.698	1.00 40.95	Α	
ATOM	165	NZ	LYS	56	43.117	34. 537	31.685	1.00 43.33	Α	N
ATOM	166	C	LYS	56	41.240	40.844	33. 252	1.00 30.03	Α	C
ATOM	167	Ō	LYS	56	40.839	41.648	34.088	1.00 28.24	A	0
ATOM	168	N	LEU	57	41.286	41.120	31.956	1.00 30.20	A	N
ATOM	169	CA	LEU	57	40.836	42. 404	31.437	1.00 29.43	A	C
ATOM	170	CB	LEU	57	42.022	43. 233	30. 934	1.00 30.04	A	C .
ATOM	171	CG	LEU	57	43. 230	43. 474	31.844	1.00 32.13	A	C
ATOM	172		LEU	57	44. 123	44. 524	31.194	1.00 29.05	A	· C
ATOM	173		LEU	57	42.777	43. 949	33. 230	1.00 34.11	A	C C
ATOM	174	C	LEU	57 57	39.911	42. 132	30. 271	1.00 28.16	A	0
ATOM	175	0	LEU	57	39.668	40. 980	29. 914 29. 676	1.00 28.60 1.00 26.69	, A , A	N N
ATOM	176	N	TYR	58	39. 394	43. 196 43. 050	28. 518	1.00 25.82	A	Č
ATOM	177	CA	TYR	58 50	38. 530 37. 071	42. 890	28. 934	1.00 25.52	A	č
ATOM	178 179	CB CG	TYR TYR	58 58	36. 195	42. 420	27. 797	1.00 26.86	Ä	č
ATOM	180	CD1		58	36. 051	41. 062		1.00 26.92	Ä	Č
MOTA	181		TYR	58	35. 294		26. 429	1.00 26.28	A	Ç,
ATOM ATOM	182		TYR	58	35. 557		26. 965	1.00 25.26	A	Č
ATOM	183		TYR	58	34. 803		25. 882	1.00 26.13	A	C
ATOM	184	CZ	TYR	58	34. 675		25.619	1.00 25.74	Α	С
ATOM	185	ОH	TYR	58	33. 928		24. 541	1.00 27.32	Α	0 ·
ATOM	186	C	TYR	58	38. 681	44. 288	27.647	1.00 24.95	Α	C
ATOM	187	Ō	TYR	58	37.837		27.680	1.00 24.68	Α	0
ATOM	188	Ň	SER	59	39.763	44. 338	26.876	1.00 24.05	A	Ŋ
ATOM	189	CA	SER	59	40.037	45.470	25.997	1.00 24.31	A	C
ATOM	190	CB	SER	59	41.547		25.817	1.00 24.38	A	C
ATOM	191	0G	SER	59	42.187		27. 051	1.00 28.99	· A	0
ATOM	192	C	SER	59	39.405				A	C
ATOM	193	0	SER	59	39. 795				A	0
ATOM	194		LEU	60	38. 430				A	N
ATOM	195	CA	LEU	60	37. 765	46.073	23.031	1.00 22.96	Α	C

			FIG. 4-5	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	196 CB LEU 197 CG LEU 198 CD1 LEU 199 CD2 LEU 200 C LEU 201 0 LEU 202 N ARG 203 CA ARG 204 CB ARG 205 CG ARG 206 CD ARG 207 NE ARG 208 CZ ARG 209 NH1 ARG 210 NH2 ARG 211 C ARG 211 C ARG 212 O ARG 213 N TRP 214 CA TRP 215 CB TRP 216 CG TRP 217 CD2 TRP 218 CE2 TRP 218 CE2 TRP 219 CE3 TRP 219 CE3 TRP 210 CD1 TRP 221 NE1 TRP 222 CZ2 TRP 223 CZ3 TRP 224 CH2 TRP 225 C TRP 226 O TRP	60 60 60 60 61 61 61 61 61 61 62 62 62 62 62 62 62 62 62 62	36. 256 45. 910 23. 228 1. 00 21. 27 A 35. 528 46. 977 24. 048 1. 00 20. 80 A 35. 373 48. 227 23. 208 1. 00 19. 95 A 34. 159 46. 466 24. 488 1. 00 18. 91 A 38. 072 47. 356 22. 279 1. 00 23. 42 A 38. 507 48. 340 22. 869 1. 00 23. 10 A 37. 862 47. 339 20. 971 1. 00 25. 94 A 38. 102 48. 522 20. 153 1. 00 27. 08 A 39. 364 48. 323 19. 299 1. 00 29. 17 A 40. 545 47. 713 20. 076 1. 00 34. 91 A 41. 790 48. 612 20. 088 1. 00 38. 62 A 42. 423 48. 715 18. 772 1. 00 41. 78 A 43. 37 47. 871 18. 299 1. 00 41. 78 A 43. 821 48. 042 17. 076 1. 00 43	A C C C A C C C A C C C A C C C A C A C C A C A C C A C A C C A C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	226 O TRP 227 N ILE 228 CA ILE 229 CB ILE 230 CG2 ILE 231 CG1 ILE 232 CD1 ILE 233 C ILE 234 O ILE 235 N SER 236 CA SER 237 CB SER 236 CA SER 237 CB SER 238 OG SER 239 C SER 240 O SER 241 N ASP 242 CA ASP 243 CB ASP 244 CG ASP	62 63 63 63 63 63 63 64 64 64 65 65 65	35. 241	A O N C C A C C C A C C C C A C

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										(Continued)
		•			FI	G. 4	- 6			
ATOM	245	0D1	ASP	65	27.070	54. 235	18.465	1.00 43.44	A	0
ATOM	246	0D2	ASP	65	25. 281	53.986	17. 211	1.00 44.76	A	0
ATOM	247	C	ASP	65	27.640	51.064	17.128	1.00 34.55	A	C
ATOM	248	0	ASP	65	26.753	51.091	17. 981	1.00 33.76	A	0
ATOM	249	N	HIS	66	28. 023	49.946	16.520	1.00 34.31	A	N
ATOM	250	CA	HIS	66	27. 369	48.679	16.807	1.00 35.30	A	C
ATOM	251	CB	HIS	66	26. 555	48. 229	15.589	1.00 37.74	A	C
ATOM	252	CG	HIS	66	25. 648	49. 288	15.052	1.00 42.72	A A	C
ATOM	253		HIS	66 cc	24. 298	49. 393	15.056 14.455	1.00 44.80 1.00 45.16	A	N
ATOM	254		HIS	66 cc	26. 121	50. 438 51. 206	14.433	1.00 45.10	Ä	C
ATOM	255		HIS	66 66	25. 101 23. 984	51. 200 50. 595	14.114	1.00 46.24	A	Ň
ATOM	256	NEZ C	HIS HIS	66 66	28. 314	47. 555	17. 223	1.00 33.78	Ä	Ċ
ATOM ATOM	257 258	0	HIS	66	27. 966	46. 736	18.068	1.00 34.67	A	Ö
ATOM	259	N	GLU	67	29. 502	47. 501	16.635	1.00 31.93	Ā	N
ATOM	260	CA	GLU	67	30. 432	46. 434	16.979	1.00 31.45	A	C
ATOM	261	CB	GLU	67	30. 557	45. 463	15.801	1.00 31.46	Α	
ATOM	262	CG	GLU	67	30.356	46.103	14.447	1.00 33.17	Α	C C C
ATOM	263	CD	GLU	67	30.357	45.092	13.311	1.00 35.48	Α	
ATOM	264		GLU	67	29.607	44.090	13. 394	1.00 32.44	Α	0
ATOM	265	0E2	GLU	67	31.104	45.306	12.329	1.00 36.60	A	0
ATOM	266	C	GLU	67	31.818	46.866	17.442	1.00 29.97	A	C
ATOM	267	0	GLU	67	32. 240	48.003	17. 241	1.00 30.44	A	0
ATOM	268	N	TYR	68	32.513	45.940	18.088	1.00 29.07	A	N
ATOM	269	CA	TYR	68	33. 863	46. 190	18.567	1.00 28.87	A	C
ATOM	270	CB	TYR	68	33.866	46. 447	20.073	1.00 26.31	A	C C
ATOM	271	CC	TYR	68	33.307	45. 324 45. 376	20. 917 21. 400	1.00 23.19 1.00 21.93	A A	C
ATOM	272	CD1	TYR TYR	68 69	32.000 31.497	43. 370	21.400	1.00 21.33	A	C
MOTA MOTA	273 274		TYR	68 68	34. 102	44. 232	21. 281	1. 00 23. 23	A	č
ATOM	275		TYR	68	33. 610	43. 225	22. 110	1.00 22.67	A	č
ATOM	276	CZ	TYR	68	32. 304	43. 305	22. 582	1.00 22.02	Ä	Č.
ATOM	277	OH	TYR	68	31.810	42. 321	23. 403	1.00 22.72	Ā	0
ATOM	278	C	TYR	68	34. 747		18.256	1.00 29.51	Α	C
ATOM	279	ŏ	TYR	68	34. 244	43.885	18.028	1.00 28.32	Α	0
ATOM	280	Ň	LEU	69	36.058	45. 202	18.233	1.00 29.87	Α	N
ATOM	281	CA	LEU	69	36.986	44.115	17.963	1.00 32.20	Α	С
ATOM	282	CB	LEU	69	38. 154	44.602	17. 106	1.00 30.73	A	C
ATOM	283	CG	LEU	69	37. 761	45.065	15. 700	1.00 30.62	A	C
ATOM	284		LEU	69	38. 978	45.629	14. 963	1.00 29.98	A	C
ATOM	285		LEU	69	37. 164		14. 943	1.00 30.17	A	C
ATOM	286	C	LEU	69	37. 492	43. 588	19. 292	1.00 34.73	A	C
ATOM	287	0	LEU	69 70	37. 474		20. 294	1.00 34.80	A	0 N
ATOM	288	N	TYR	70	37. 927	42.334	19.305	1.00 37.39 1.00 42.16	A	N
ATOM	289	CA	TYR	70 70	38. 423	41.726	20.528	1.00 42.10	A A	C C
ATOM	290	CB	TYR	70 70	37. 251	41.359	21.444 22.799	1.00 42.00	A	C
ATOM	291	CG	TYR TYR	70 70	37. 689 38. 400	40.866 41.697	23.657	1.00 43.00	A	Č
ATOM ATOM	292 293		TYR	70 70	38. 837		24. 892	1.00 44.69	A	č
UIOM	400	נוטי		10	00.001	11. 500				=

(Continued) FIG. 4-7 C ATOM 294 CD2 TYR 70 37.421 39.563 23. 213 1.00 43.93 Α C **ATOM** 295 CE2 TYR 70 37.853 39.104 24.452 1.00 44.83 Α CZTYR 70 38.563 39.959 25.286 1.00 45.17 C 296 Α ATOM 39.004 39.532 26.516 0 ATOM 297 0HTYR 70 1.00 47.21 Α 70 39.249 40.480 20.240 1.00 45.46 A C **ATOM** 298 C TYR **ATOM** 299 0 TYR 70 38.976 39.752 19.287 1.00 46.31 A 0 LYS 40.254 40.231 21.072 ATOM 300 N 71 1,00 49,93 A N CA LYS 41.113 39.064 20.895 1.00 54.71 A C **ATOM** 301 71 LYS C CB 42.580 39.460 21.054 1.00 54.14 Α ATOM 302 71 C **ATOM** CG LYS 43.075 40.455 20.031 1.00 56.37 Α 303 71 CD ·LYS 44.559 40.712 20.226 C ATOM 304 71 1.00 58.61 Α C **ATOM** CE LYS 45.126 41.628 19.159 1.00 58.78 A 305 71 LYS 46.590 **ATOM** 306 NZ 71 41.830 19.361 1.00 60.82 A N **ATOM** C LYS 40.790 37.952 21.889 1.00 57.38 C 307 71 Α **ATOM** 0 LYS 41.109 38.062 23.075 308 1.00 58.38 0 71 Α N **ATOM** 309 GLN 40.158 36.884 21.406 1.00 60.30 72 A N CA **ATOM** 310 GLN 72 39.816 35.750 22.261 1.00 63.23 A C **ATOM** 311 CB GLN 72 38.902 34.775 21.526 1.00 64.07 C A 38.313 312 CG GLN 72 33.695 22.417 **ATOM** 1.00 65.84 Α C **ATOM** 313 CD GLN 72 37.270 34.240 23.375 1.00 66.33 C Α ATOM 72 36.251 34.790 314 OE1 GLN 22.952 1.00 67.19 0 Α NE2 GLN 37.519 34.092 ATOM 315 72 24.671 1.00 66.80 A N 41. 122 41. 563 22.607 **ATOM** 316 C GLN 72 35.049 1.00 65.34 A C 0 ATOM 317 GLN 72 35.058 23.760 1.00 67.00 0 A **ATOM** N GLU 73 41.736 34.442 21.597 318 1.00 66.09 Α. N CA GLU **ATOM** 319 73 43.012 33.763 21.775 1.00 67.12 Α C **ATOM** 320 CB GLU 73 43.008 32.420 21.046 1.00 68.53 C Α CG **ATOM** 321 GLU 73 41.974 31.433 21.570 1.00 71.35 C A **ATOM** 322 CD GLU 73 42.223 31.026 23.012 1.00 72.71 C Α OE1 GLU ATOM 323 73 41.491 30.147 23.517 1.00 73.51 0 A **ATOM** 324 OE2 GLU 73 43.147 31.585 23.643 1.00 74.16 A , **0 ATOM** 325 C **GLU** 73 44.076 34.681 21.184 1.00 66.83 C A **ATOM** 326 0 **GLU** 73 35, 592 44.563 21.857 1.00 67.65 . 0 Α N ATOM 327 ASN 74 44.430 34.442 19.924 1.00 65.38 A N **ATOM** 328 CA ASN 45.411 35.273 19.236 74 1.00 63.38 C A CB **ATOM** 329 ASN 74 46.661 34.466 18.889 1.00 64.38 A C **ATOM** 330 CG ASN 74 47.654 34.422 20.034 1.00 66.10 A C **ATOM** 331 OD1 ASN 48.128 35.463 20.496 74 1.00 65.51 A 0 47. 973 332 20.503 ATOM ND2 ASN 33.216 74 1.00 66.62 A N **ATOM** 333 C ASN 44.794 35.859 74 17.977 1.00 61.55 C A 334 0 **ATOM** ASN 74 45.384 36.714 17.318 1.00 62.15 A 0 **ATOM** 335 N ASN 75 43.597 35.390 17.647 1.00 58.67 A N ATÓM 336 CA ASN 75 42.888 35.886 16.481 1.00 55.82 C A C **ATOM** 337 CB ASN 75 42.023 34.785 15.871 1.00 57.81 Α CG 1.00 58.63 C ATOM 338 ASN 75 41.410 33.887 16.916 Α **ATOM** 339 OD1 ASN 34.358 17.909 1.00 59.69 0 75 40.857 Α ATOM 340 ND2 ASN 75 41.500 32.580 16.697 1.00 58.92 A N 341 ASN 16.918 1.00 52.82 C ATOM C 75 42.017 37.045 A

SUBSTITUTE SHEET (RULE 26)

41.630

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ASN

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ATOM

			FIG. 4-8	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	343 N ILE 344 CA ILE 345 CB ILE 346 CG2 ILE 347 CG1 ILE 348 CD1 ILE 349 C ILE 350 O ILE 351 N LEU 352 CA LEU 355 CD1 LEU 355 CD1 LEU 356 CD2 LEU 357 C LEU 357 C LEU 358 O LEU 359 N VAL 361 CB VAL 361 CB VAL 362 CG1 VAL 363 CG2 VAL 364 C VAL 365 O VAL 366 N PHE 367 CA PHE 368 CB PHE 369 CG PHE 368 CB PHE 369 CG PHE 370 CD1 PHE 371 CD2 PHE 372 CE1 PHE 373 CE2 PHE 374 CZ PHE 375 C PHE 377 N ASN 378 CA ASN 379 CB ASN	76 76 76 76 76 77 77 77 77 77 77 78 78 78 79 79 79 79 79 79 79 79 79 79 80 80 80 80 80	41. 715	(Continued) A
ATOM ATOM ATOM ATOM	382 ND2 ASN 383 C ASN 384 O ASN 385 N ALA	80 80 80 80 81	24. 979 42. 587 18. 182 1. 00 26. 87 24. 980 42. 866 15. 974 1. 00 26. 94 27. 405 42. 874 19. 289 1. 00 28. 06 26. 991 42. 024 20. 066 1. 00 28. 61 27. 566 44. 140 19. 642 1. 00 28. 12 27. 250 44. 579 20. 991 1. 00 29. 16	A N A C A O A N A C
ATOM ATOM ATOM ATOM ATOM ATOM	386 CA ALA 387 CB ALA 388 C ALA 389 O ALA 390 N GLU 391 CA GLU	81 81 81 82 82	27. 250 44. 373 20. 331 1. 00 23. 10 27. 503 46. 075 21. 119 1. 00 27. 93 25. 818 44. 254 21. 413 1. 00 31. 04 25. 582 43. 769 22. 527 1. 00 30. 16 24. 870 44. 506 20. 516 1. 00 32. 39 23. 461 44. 282 20. 809 1. 00 34. 46	A C A C A O A N A C

					FI	G. 4	- 9			(Continued)
ATOM ATOM	392 393	CB CG	GLU GLU	82 82	22. 602 21. 115	44. 794 44. 827	19.655 19.968	1.00 36.97 1.00 40.49	A A	C C
ATOM	394	CD	GLU	82	20. 313	45. 538	18.894	1.00 44.05	A	č
ATOM	395	0E1		82	20. 343	45.087	17.726	1.00 45.13	Ā	0
ATOM	396	0E2		82	19.652	46.551	19.220	1.00 45.61	A	0
ATOM	397	C	GLU	82	23.042	42.853	21.153	1.00 33.95	A	C
ATOM	398	0	GLU	82	22.055	42.662	21.864	1.00 32.29	Α	0
ATOM	399	N	TYR	83	23. 777	41.857	20.666	1.00 33.23	A	N
ATOM	400	CA	TYR	83	23. 423	40.468	20.947	1.00 33.39	A	Č
ATOM	401	CB	TYR	83	22.846	39. 810	19.686	1.00 34.54	A	C
ATOM	402	CG	TYR	83	21.690	40.594	19.109	1.00 34.80	A	C
ATOM	403	CD1	TYR	83	20. 558	40.859	19.878	1.00 35.22	A	C
ATOM	404	CE1	TYR	83	19. 527	41.657	19.396	1.00 36.27	A	C
ATOM	405		TYR	83	21. 759	41.139	17.828	1.00 35.71	A	C
ATOM ATOM	406 407	CE2 CZ	TYR TYR	83	20. 731	41.940	17. 331	1.00 37.42	A	C
ATOM	408	OH	TYR	83 83	19. 619 18. 624	42. 200 43. 044	18. 125 17. 675	1.00 37.70 1.00 37.69	A A	C 0
ATOM	409	C	TYR	83	24. 582	39.644	21.494	1.00 37.09	A	Č
ATOM	410	Ö	TYR	83	24. 396	38. 511	21. 934	1.00 33.13	A	0
ATOM	411	N	GLY	84	25. 777	40. 217	21. 476	1.00 32.51	A	N
ATOM	412	CA	GLY	84	26. 933	39. 513	21. 995	1.00 33.40	A	Č
ATOM	413	C	GLY	84	27. 454	38. 395	21.114	1.00 33.92	A	č
ATOM	414	Ö	GLY	84	28. 329	37.639	21.530	1.00 33.21	Ä	Ŏ
ATOM	415	N	ASN	85	26.918	38. 269	19.904	1.00 35.26	A	N
ATOM	416	CA	ASN	85	27.388	37. 233	18.993	1.00 37.43	A	C
ATOM	417	CB	ASN	85	26.258	36.780	18.072	1.00 38.34	Α	C
ATOM	418	CG	ASN	85	25. 764	37.878	17.166	1.00 40.02	Α	C
ATOM	419	0D1	ASN	85	25.694	39.040	17.561	1.00 39.96	A	0
ATOM	420		ASN	85	25. 394	37.496	15.950	1.00 41.91	A	N
ATOM	421	C	ASN	85	28. 556	37. 794	18. 188	1.00 38.80	A	C
ATOM	422	0	ASN	85	28. 687	39.011	18.035	1.00 40.05	A	0
ATOM	423	N	SER	86	29. 410	36.920	17.670	1.00 39.14	A	N
ATOM ATOM	424 425	CA ⁻ CB	SER	86	30. 565	37. 393	16.926	1.00 39.30	A	C
ATOM	425 426	OG	SER SER	86 86	31.723	37.587	17.895	1.00 38.90	A	C
ATOM	427	C	SER	86	32. 041 31. 023	36. 356 36. 482	18. 515 15. 798	1.00 35.77 1.00 39.94	A	0
ATOM	428	0	SER	86	30. 287	35. 622	15. 323	1.00 39.94	A A	C 0
ATOM	429	N	SER	87	32. 264	36. 701	15. 382	1.00 40.59	A	N
ATOM	430	CA	SER	87	32. 916	35. 929	14. 333	1.00 40.98	Ä	Č
ATOM	431	CB	SER	87	32. 152	36.053	13.010	1.00 39.16	A	č
ATOM	432	0G	SER	87	31.727	37. 376	12. 789	1.00 39.90	A	ŏ
ATOM	433	Č	SER	87	34. 353	36. 433	14. 194	1.00 41.10	Ä	č
ATOM	434	Ŏ	SER	87	34. 691	37. 517	14. 682	1.00 41.07	Ä	ŏ
ATOM	435	N	VAL	88	35. 206	35.646	13.548	1.00 41.07	Ä.	Ň
ATOM	436	CA	VAL	88	36.596	36.043	13.402	1.00 41.43	A	Ċ
ATOM	437	CB	VAL	88	37.502	34.836	13.114	1.00 41.29	Α	С
ATOM	438	CG1	VAL	88	38.949	35. 295	13.013	1.00 41.30	Α	С
ATOM	439		VAL	88	37.361	33.808	14. 222	1.00 40.28	Α	C
ATOM	440	C	VAL	88	36. 827	37.096	12. 331	1.00 41.63	Α	C

				(Continued)
	. •		FIG. 4-10	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	441 O VAL 442 N PHE 443 CA PHE 444 CB PHE 445 CG PHE 446 CD1 PHE 447 CD2 PHE 448 CE1 PHE 449 CE2 PHE 450 CZ PHE 451 C PHE 452 O PHE 452 O PHE 453 N LEU 454 CA LEU 455 CB LEU 456 CG LEU 457 CD1 LEU 458 CD2 LEU 458 CD2 LEU 459 C LEU 459 C LEU 460 O LEU 461 N GLU 462 CA GLU 463 CB GLU 463 CB GLU 464 CG GLU 465 CD GLU 466 OE1 GLU 467 OE2 GLU 468 C GLU 467 OE2 GLU 468 C GLU 467 OE2 GLU 467 OE2 GLU 468 C GLU 467 OE2 GLU 467 OE2 GLU 468 C GLU 467 OE2 GLU 467 OE2 GLU 468 C GLU 469 O GLU 470 N ASN 471 CA ASN 472 CB ASN 474 OD1 ASN 475 ND2 ASN 476 C ASN 477 O ASN 478 N SER 479 CA SER 480 CB SER 481 OG SER 481 OG SER	88 89 89 89 89 89 89 89 89 89	36. 548 36. 885 11. 154 1. 00 41. 38 37. 343 38. 238 12. 767 1. 00 42. 23 37. 641 39. 347 11. 880 1. 00 42. 51 37. 769 40. 637 12. 699 1. 00 40. 84 37. 990 41. 865 11. 870 1. 00 39. 96 39. 217 42. 103 11. 265 1. 00 39. 62 36. 963 42. 778 11. 678 1. 00 40. 08 39. 415 43. 231 10. 480 1. 00 39. 60 37. 154 43. 911 10. 894 1. 00 39. 50 38. 956 39. 021 11. 186 1. 00 43. 57 39. 156 39. 335 10. 019 1. 00 45. 92 41. 143 38. 001 11. 380 1. 00 48. 60 42. 071 39. 213 11. 366 1. 00 48. 66 43. 033 39. 305 10. 184 1. 00 49. 47 42. 236 39. 408 8. 889 1. 00 50. 17 43. 933 40. 515 10. 346 1. 00 50. 91 41. 718 36. 907 12. 267 1. 00 50. 84 42. 063 37. 159 <t< td=""><td>(Continued) A</td></t<>	(Continued) A
ATOM ATOM ATOM ATOM ATOM ATOM	482 C SER 483 O SER 484 N THR 485 CA THR 486 CB THR 487 OG1 THR	93 93 94 94 94	47. 296 33. 030 9. 853 1. 00 62. 23 48. 314 32. 765 9. 213 1. 00 62. 82 46. 552 34. 103 9. 618 1. 00 62. 37 46. 852 35. 036 8. 541 1. 00 62. 69 45. 982 36. 298 8. 659 1. 00 63. 25 46. 469 37. 302 7. 759 1. 00 63. 59	A 0 A N A C A C A 0
ATOM ATOM	488 CG2 THR 489 C THR	94 94	46.003 36.821 10.080 1.00 64.14 48.306 35.464 8.377 1.00 62.28	A C A C

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			FIC 4-11	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	490 O TH 491 N PH 492 CA PH 493 CB PH 494 CG PH 495 CD1 PH 496 CD2 PH 497 CE1 PH 498 CE2 PH 499 CZ PH 500 C PH 501 O PH	E 95 95 95 95 95 95 95 95 95 95 95 95 95	FIG. 4 - 11 48. 882 35. 295 7. 303 1. 00 61. 92 48. 908 36. 013 9. 426 1. 00 62. 57 50. 290 36. 473 9. 322 1. 00 63. 04 50. 414 37. 889 9. 897 1. 00 61. 98 49. 456 38. 869 9. 289 1. 00 61. 01 48. 248 39. 155 9. 911 1. 00 60. 97 49. 742 39. 473 8. 073 1. 00 60. 73 47. 337 40. 026 9. 330 1. 00 60. 46 48. 838 40. 343 7. 483 1. 00 60. 09 47. 633 40. 621 8. 113 1. 00 61. 07 51. 346 35. 571 9. 956 1. 00 63. 20 52. 178 36. 035 10. 736 1. 00 63. 66 51. 283 24. 388 9. 611 1. 00 63. 27	A O A N A C A C A C A C A C A C A C A C A C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	502 N AS 503 CA AS 504 CB AS 505 CG AS 506 OD1 AS 507 OD2 AS 508 C AS 509 O AS 510 N GL 511 CA GL 512 CB GL	P 96 P 96 P 96 P 96 P 96 P 96 U 97 U 97 U 97	51. 323 34. 288 9. 611 1. 00 63. 37 52. 298 33. 347 10. 149 1. 00 64. 05 51. 771 31. 913 10. 044 1. 00 65. 11 50. 747 31. 589 11. 115 1. 00 65. 73 49. 758 32. 342 11. 240 1. 00 66. 41 50. 929 30. 580 11. 829 1. 00 65. 32 53. 621 33. 470 9. 399 1. 00 63. 82 54. 696 33. 433 10. 001 1. 00 64. 05 53. 540 33. 619 8. 083 1. 00 62. 95 54. 740 33. 754 7. 271 1. 00 62. 73 54. 596 32. 964 5. 965 1. 00 65. 91 54. 954 31. 478 6. 064 1. 00 68. 84	A N A C A C A C A O A O A C A O A C A O A C A C A C A C A C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	514 CD GL 515 OE1 GL 516 OE2 GL 517 C GL 518 O GL 519 N PH 520 CA PH 521 CB PH 522 CG PH 523 CD1 PH 524 CD2 PH 525 CE1 PH	U 97 U 97 U 97 U 97 E 98 E 98 E 98 E 98	53. 945 30. 657 6. 850 1. 00 70. 64 54. 160 29. 432 6. 988 1. 00 71. 38 52. 939 31. 228 7. 325 1. 00 71. 80 55. 039 35. 220 6. 963 1. 00 60. 82 55. 462 35. 557 5. 857 1. 00 60. 31 54. 818 36. 084 7. 952 1. 00 58. 68 55. 067 37. 513 7. 797 1. 00 55. 93 54. 200 38. 319 8. 765 1. 00 55. 47 54. 272 39. 801 8. 542 1. 00 54. 84 53. 712 40. 372 7. 404 1. 00 53. 07 54. 931 40. 624 9. 450 1. 00 53. 89 53. 808 41. 743 7. 173 1. 00 53. 28	A C A O A C A C A C A C A C A C A C A C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	526 CE2 PH 527 CZ PH 528 C PH 529 O PH 530 N GL 531 CA GL 532 C GL 533 O GL 534 N HI 535 CA HI 536 CB HI 537 CG HI 538 CD2 HI	E 98 E 98 Y 99 Y 99 Y 99 S 100 S 100 S 100	55. 032 41. 997 9. 226 1. 00 53. 18 54. 470 42. 556 8. 087 1. 00 52. 22 56. 536 37. 820 8. 060 1. 00 54. 61 57. 041 38. 878 7. 686 1. 00 53. 80 57. 215 36. 885 8. 713 1. 00 53. 53 58. 624 37. 061 9. 004 1. 00 52. 08 58. 908 38. 188 9. 972 1. 00 51. 18 60. 037 38. 673 10. 051 1. 00 51. 30 57. 884 38. 607 10. 706 1. 00 50. 21 58. 026 39. 681 11. 686 1. 00 49. 15 57. 810 41. 049 11. 028 1. 00 48. 84 58. 850 41. 410 10. 014 1. 00 49. 22 58. 759 41. 613 8. 679 1. 00 49. 42	A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C A C C C A C

٠.					EI	G. 4	- 1 9			(Continued)	
ATOM	539	ND1		100	60.170	41.627	10.346	1.00 49.70	A	N C	
ATOM	540	CE1		100	60.848	41.951	9. 259	1.00 49.10	A	C	
ATOM . ATOM	541 542	NE2 C	HIS	100 100	60. 015 57. 011	41.949 39.511	8. 234 12. 810	1.00 50.14 1.00 48.06	A A	N C	
ATOM	543	0	HIS	100	55. 920	38. 977	12.602	1.00 48.00	A	0	
ATOM	544	N	SER	101	57. 377	39. 958	14. 005	1.00 46.66	A	N	
ATOM	545	CA	SER	101	56. 467	39.878	15. 136	1.00 45.88	A	Č	
ATOM	546	CB	SER	101	57. 247	39. 802	16. 446	1.00 47.41	A	č	
ATOM	547	OG	SER	101	58. 118	38. 685	16. 447	1.00 51.04	Ä	Ö	
ATOM	548	Č	SER	101	55.617	41.142	15. 112	1.00 44.53	A	Č	
ATOM	549	Ō	SER	101	56.133	42.248	15. 282	1.00 44.41	A	0 .	
ATOM	550	N	ILE	102	54.319	40.976	14.877	1.00 41.90	Α	N	
ATOM	551	CA	ILE	102	53.409	42.109	14.833	1.00 38.95	Α	С	
ATOM	552	CB	ILE	102	52.106	41.732	14. 117	1.00 38.54	Α	C C	
ATOM	55 3	CG2		102	51.153	42, 926	14. 103	1.00 38.18	Α	С	
ATOM	554	CG1		102	52.424	41.288	12.686	1.00 37.65	A	C	
ATOM	555	CD1	ILE	102	51.243	40. 733	11. 937	1.00 37.11	A	C	
ATOM	556	C	ILE	102	53. 104	42.597	16. 244	1.00 38.00	A	C	
ATOM	557	0	ILE	102	52. 441	41.919	17. 024	1.00 38.06	A	0	
ATOM	558	N	ASN	103	53. 601	43. 787	16.556	1.00 37.54	A	N	
ATOM	559	CA	ASN	103	53. 429	44. 399	17. 867	1.00 36.65	A	C	
ATOM ATOM	560 561	CB CG	ASN ASN	103 103	54. 437 54. 219	45. 530 46. 308	18. 039 19. 315	1.00 37.69 1.00 39.56	A A	C C	
ATOM	562	OD1	ASN	103	54. 655	45. 891	20. 388	1.00 39.00	A	0	
ATOM	563		ASN	103	53. 528	47. 439	19. 211	1.00 43.00	A	N	
ATOM	5 6 4	C	ASN	103	52. 031	44. 953	18. 116	1.00 35.79	A	C	
ATOM	565	ŏ	ASN	103	51.532	44.910	19. 237	1.00 35.79	A	ŏ ·	
ATOM	566	Ň	ASP	104	51.405	45.490	17. 078	1.00 34.43	Ä	Ň	
ATOM	567	CA	ASP	104	50.079	46.067	17. 236	1.00 33.27	Ä	Č	
ATOM	568	CB	ASP	104	50. 200	47.388	17.998	1.00 34.38	Α	C	
ATOM	569	CG	ASP	104	48.896	47.823	18.618	1.00 34.79	Α	C	
ATOM	570			104	48.916	48.699	19.509	1.00 33.92	Α	0	
ATOM	571		ASP	104	47.852	47. 289	18. 207	1.00 36.80	Α	0	
ATOM	572	C	ASP	104	49. 436	46. 281	15. 865	1.00 32.32	A	Č	
ATOM	573	0	ASP	104	50. 124	46.326	14.850	1.00 32.03	A	0	
ATOM	574	N	TYR	105	48. 118	46.405	15. 834	1.00 31.15	A	N	
ATOM	575	CA	TYR	105	47. 421	46.580	14. 570	1.00 32.24	A	C	
ATOM	576	CB	TYR	105	46.672	45. 296	14. 223	1.00 34.70	A	C	
ATOM ATOM	577 578	CG CD1	TYR TYR	105	45. 443	45.088 45.636	15.072	1.00 37.73	A	C C	
ATOM	579		TYR	105 105	44. 220 43. 098	45. 510	14. 698 15. 506	1.00 37.51 1.00 40.43	A A	C	
ATOM	580		TYR	105	45. 514	44. 395	16. 284	1.00 40.43	A	C	
ATOM	581	CE2		105	44. 393	44. 263	17. 103	1.00 40.75	A	Č	
ATOM	582	CZ	TYR	105	43. 191	44. 829	16.705	1.00 41.19	A	č	
ATOM	583	OH	TYR	105	42. 088	44. 755	17. 519	1.00 44.27	Ä	ŏ ·	
ATOM	584	C	TYR	105	46. 441	47. 743	14. 638	1.00 31.43	Ä	č	
ATOM	585	Ŏ	TYR	105	46. 133	48. 249	15.715	1.00 30.78	A	Ŏ	
ATOM	586	Ň	SER	106	45. 940	48. 152	13.479	1.00 30.16	Ä	N	
ATOM	587	CA	SER	106	45.000	49.261	13.415	1.00 29.23	Α	C	

		म	`IG. 4	- 13			(Continued)
ATOMATOMATOMATOMATOMATOMATOMATOMATOMATOM	588 CB SER 589 OG SER 590 C SER 591 O SER 592 N ILE 593 CA ILE 594 CB ILE 595 CG2 ILE 596 CG1 ILE 597 CD1 ILE 598 C ILE	106 45.3 106 44.9 106 44.6 107 42.8 107 41.9 107 40.6 107 39.8 107 40.9 107 41.9	924 51.668 146 49.187 657 49.085 335 49.240 922 49.171 648 48.352 657 48.620 970 46.859 980 46.457	13. 457 13. 090 12. 157 11. 051 12. 331 11. 198 11. 544 10. 522 11. 551 12. 568 10. 743	1. 00 29. 81 1. 00 32. 32 1. 00 27. 65 1. 00 28. 57 1. 00 28. 07 1. 00 27. 70 1. 00 25. 83 1. 00 26. 35 1. 00 25. 36 1. 00 23. 77 1. 00 26. 85	A A A A A A A A	C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	599 O ILE 600 N SER 601 CA SER 602 CB SER 603 OG SER 604 C SER 605 O SER 606 N PRO 607 CD PRO 608 CA PRO 609 CB PRO	107 41. 1 108 41. 5 108 41. 1 108 41. 3 108 40. 4 108 39. 6 108 38. 8 109 39. 2 109 40. 0 109 37. 8 109 37. 8	178 51. 420 507 50. 757 13 52. 035 331 52. 033 158 51. 119 339 52. 253 357 51. 310 241 53. 506 025 54. 751 339 53. 794	11. 557 9. 432 8. 862 7. 346 6. 700 9. 169 9. 206 9. 393 9. 302 9. 693 9. 439	1. 00 26. 55 1. 00 27. 57 1. 00 26. 94 1. 00 26. 30 1. 00 23. 63 1. 00 27. 22 1. 00 26. 49 1. 00 28. 50 1. 00 29. 19 1. 00 29. 39 1. 00 30. 19	A A A A A A A A	0 N C C O C O N C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	610 CG PRO 611 C PRO 612 O PRO 613 N ASP 614 CA ASP 615 CB ASP 616 CG ASP 617 OD1 ASP 618 OD2 ASP 619 C ASP	109 39. 0 109 36. 8 109 35. 9 110 37. 0 110 36. 1 110 36. 2 110 38. 2 110 38. 0 110 36. 2	55. 775 542 52. 993 501 52. 425 546 52. 935 520 52. 202 541 52. 673 513 52. 432 526 51. 397 53. 274	9. 899 8. 852 9. 391 7. 540 6. 676 5. 226	1. 00 28. 76 1. 00 29. 21 1. 00 30. 65 1. 00 29. 41 1. 00 28. 98 1. 00 27. 99 1. 00 27. 91 1. 00 28. 41 1. 00 29. 14 1. 00 29. 06	A A A A A A A	C C O N C C C O
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	620 O ASP 621 N GLY 622 CA GLY 623 C GLY 624 O GLY 625 N GLN 626 CA GLN 627 CB GLN 628 CG GLN	110 35.6 111 37.3 111 37.8 111 37.8 111 37.8 112 38.4 112 38.5 112 38.7 112 37.3	335 49. 971 48 50. 196 349 48. 766 390 48. 064 356 46. 837 405 48. 818 346 48. 217 477 49. 171 336 49. 442	5. 953 7. 589 7. 702 6. 470 6. 402 5. 503 4. 287 3. 109 2. 749	1.00 30.84 1.00 28.25 1.00 28.14 1.00 29.53 1.00 31.16 1.00 29.61 1.00 29.74 1.00 29.94 1.00 31.79	A A A A A A A	C O N C C O N C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	629 CD GLN 630 OE1 GLN 631 NE2 GLN 632 C GLN 633 O GLN 634 N PHE 635 CA PHE 636 CB PHE	112 37.1 112 36.0 112 38.3 112 40.4 112 40.8 113 41.1 113 42.5 113 43.4	91 50. 234 975 50. 474 814 50. 644 815 47. 813 888 46. 971 41 48. 418 851 48. 106	1. 465 1. 004 0. 880 4. 390 3. 631 5. 320 5. 486 4. 900	1.00 33.24 1.00 36.27 1.00 31.73 1.00 30.31 1.00 31.75 1.00 29.82 1.00 28.23 1.00 24.48	A A A A A A	C O N C O N C C

ATOM 637 CG PHE 113 42.164 60.301 3.052 1.00 20.33 A C ATOM 638 CD1 PHE 113 44.010 48.880 2.495 1.00 20.33 A C ATOM 640 CE1 PHE 113 44.950 50.552 1.709 1.00 20.79 A C ATOM 641 CE2 PHE 113 43.805 49.121 1.150 1.00 22.78 A C ATOM 641 CE2 PHE 113 42.919 47.974 6.947 1.00 22.17 A C ATOM 642 CZ PHE 113 42.214 48.511 7.827 1.00 30.03 3. A C ATOM 644 CP PHE 113 42.214 48.511 7.827 1.00 30.31 A C ATOM 645 N 1LE 114 44.521 47.092 8.542 1.00 29.79 A O ATOM 646 CA LE 114 44.521 47.092 8.542 1.00 30.73 A C ATOM 646 CA LE 114 44.521 47.092 8.542 1.00 30.73 A C ATOM 649 CG ILE 114 48.04 44.633 8.042 1.00 30.33 A C ATOM 649 CG ILE 114 44.65 1.88 4.092 1.1007 1.00 30.30 A C ATOM 649 CG ILE 114 44.60 44.638 8.042 1.00 30.30 A C ATOM 650 DI ILE 114 45.128 45.75 10.381 1.00 32.62 A C ATOM 651 C ILE 114 46.504 44.633 8.042 1.00 30.36 A C ATOM 652 DI ILE 114 46.504 44.633 8.042 1.00 30.36 A C ATOM 653 N LEU 115 46.388 48.343 9.431 1.00 30.68 A N ATOM 655 CB LEU 115 49.135 50.941 10.131 1.00 30.03 5.7 A C ATOM 656 CB LEU 115 49.135 50.941 10.131 1.00 30.07 A C ATOM 657 CD ILE 114 49.688 88.343 9.433 1.00 30.77 A C ATOM 658 CD LEU 115 49.135 50.941 10.131 1.00 30.77 A C ATOM 657 CD ILEU 115 49.135 50.941 10.131 1.00 30.77 A C ATOM 658 CD LEU 115 49.135 50.941 10.131 1.00 30.77 A C ATOM 657 CD ILEU 115 49.135 50.941 10.131 1.00 30.77 A C ATOM 658 CD LEU 115 48.481 47.911 10.530 1.00 30.77 A C ATOM 657 CD ILEU 115 49.135 50.941 10.131 1.00 30.77 A C ATOM 658 CD LEU 115 48.388 48.343 1.00 30.77 A C ATOM 657 CD ILEU 115 48.481 47.911 10.530 1.00 30.77 A C ATOM 657 CD ILEU 115 49.135 50.941 10.131 1.00 30.77 A C ATOM 658 CD LEU 115 48.588 48.343 1.00 30.07 A C ATOM 657 CD ILEU 115 49.135 50.941 10.131 1.00 30.77 A C ATOM 658 CD LEU 115 49.848 47.188 10.048 1.00 30.77 A C ATOM 658 CD LEU 115 49.856 51.47 48.811 1.00 30.77 A C ATOM 658 CD LEU 115 49.868 51.47 48.81 1.00 30.77 A C ATOM 658 CD LEU 116 50.624 45.023 10.103 1.00 30.97 A C ATOM 658 CD LEU 116 50.664 45.023 10.00 30.90 A C ATOM 658 CD LEU 116 50.664 45.00 30.00 30.00 A C ATOM 658								(6:
ATOM 637 CG PHE 113 43.193 49.467 3.458 1.00 22.98 A C ATOM 638 CD1 PHE 113 42.164 50.301 3.052 1.00 20.83 A C ATOM 640 CE1 PHE 113 44.010 48.880 2.496 1.00 23.28 A C ATOM 640 CE1 PHE 113 41.950 50.552 1.709 1.00 20.79 A C ATOM 641 CE2 PHE 113 41.950 50.552 1.709 1.00 20.79 A C ATOM 641 CE2 PHE 113 41.950 50.552 1.709 1.00 22.78 A C ATOM 642 CZ PHE 113 42.771 49.962 0.754 1.00 22.17 A C ATOM 643 C PHE 113 42.271 49.962 0.754 1.00 22.17 A C ATOM 644 CR PHE 113 42.294 48.511 7.827 1.00 31.09 A O ATOM 645 N ILE 114 44.013 47.260 7.196 1.00 22.77 A C ATOM 645 N ILE 114 44.521 47.092 8.542 1.00 30.31 A C ATOM 645 N ILE 114 44.521 47.092 8.542 1.00 30.73 A C ATOM 646 CA ILE 114 44.521 47.092 8.542 1.00 30.73 A C ATOM 646 CC ILE 114 44.804 44.638 8.042 1.00 30.73 A C ATOM 645 N ILE 114 44.521 47.092 8.542 1.00 30.73 A C ATOM 646 CC ILE 114 44.522 47.092 8.542 1.00 30.73 A C ATOM 645 N ILE 114 44.522 47.092 8.542 1.00 30.73 A C ATOM 645 N ILE 114 46.000 47.457 8.509 1.00 30.73 A C ATOM 645 N ILE 114 46.000 47.457 8.509 1.00 30.73 A C ATOM 650 CDI ILE 114 46.000 47.457 8.509 1.00 30.59 A C ATOM 651 C ILE 114 46.000 47.457 8.509 1.00 30.59 A C ATOM 651 C ILE 114 46.000 47.457 8.509 1.00 30.59 A C ATOM 655 CB LEU 115 47.759 48.814 9.543 1.00 29.92 A C ATOM 656 CD LEU 115 49.668 51.147 7.661 1.00 28.76 A O ATOM 658 CD LEU 115 49.668 51.147 7.69 50.57 10.053 1.00 30.55 A C ATOM 656 CD LEU 115 49.668 51.147 8.814 9.543 1.00 29.92 A C ATOM 650 CD ILE 116 49.484 47.911 10.530 1.00 30.55 A C ATOM 660 N LEU 116 49.484 47.911 10.530 1.00 30.55 A C ATOM 660 N LEU 115 49.668 51.147 8.811 1.00 31.72 A C ATOM 660 N LEU 115 49.668 51.147 8.811 1.00 31.72 A C ATOM 660 N LEU 115 49.668 51.47 8.814 9.543 1.00 29.92 A C ATOM 660 N LEU 115 49.668 51.47 8.814 9.543 1.00 29.92 A C ATOM 660 N LEU 115 49.668 51.47 8.814 9.543 1.00 29.92 A C ATOM 660 N LEU 116 49.484 47.911 10.530 1.00 30.55 A C ATOM 660 N LEU 116 49.978 43.811 1.00 30.77 A C ATOM 660 N LEU 116 49.484 47.911 10.530 1.00 29.61 A C ATOM 660 N LEU 116 50.245 46.273 10.00 30.50 A C		•		EIC 1	1 /		,	(Continued)
ATOM 638 CD1 PHE 113				rig. 4	14			
ATOM 638 CD1 PHE 113	MOTA	COT CC DUE	119	12 102 10 167	2 /5Q	1 00 22 98	Α	С
ATOM 649 CED PHE 113								
ATOM 640 CE1 PHE 113 41.950 50.552 1.709 1.00 20.79 A C ATOM 641 CE2 PHE 113 43.805 49.121 1.150 1.00 22.78 A C ATOM 642 CZ PHE 113 42.919 47.974 6.947 1.00 30.31 A C ATOM 644 0 PHE 113 42.919 47.974 6.947 1.00 30.31 0 A C ATOM 644 0 PHE 113 42.234 48.511 7.827 1.00 31.09 A O ATOM 645 N ILE 114 44.013 47.260 7.196 1.00 29.70 A N ATOM 646 CA ILE 114 44.521 47.092 8.542 1.00 30.73 A C ATOM 647 CB ILE 114 44.521 47.092 8.542 1.00 30.73 A C ATOM 648 CG2 ILE 114 44.804 44.633 8.042 1.00 33.033 A C C ATOM 649 CG1 ILE 114 45.128 45.475 10.381 1.00 32.62 A C ATOM 650 CD1 ILE 114 45.128 45.475 10.381 1.00 32.62 A C ATOM 651 C ILE 114 46.000 47.457 8.509 1.00 30.59 A C ATOM 652 N ILE 115 46.384 48.343 9.423 1.00 30.68 A N ATOM 655 C ILE 115 47.769 48.814 9.543 1.00 29.92 A C ATOM 656 CG LEU 115 49.135 50.941 10.131 1.00 31.72 A C ATOM 656 CG LEU 115 49.686 51.147 8.718 1.00 33.172 A C ATOM 656 CG LEU 115 49.686 51.147 8.718 1.00 33.172 A C ATOM 656 CD LEU 115 49.686 51.147 8.718 1.00 33.77 A C ATOM 656 CD LEU 115 49.686 51.147 8.718 1.00 33.77 A C ATOM 656 CD LEU 115 49.686 51.147 8.718 1.00 33.77 A C ATOM 656 CD LEU 115 49.486 51.147 8.718 1.00 33.77 A C ATOM 656 CD LEU 115 49.486 51.147 8.718 1.00 33.77 A C ATOM 656 CD LEU 115 49.486 45.147 8.718 1.00 33.77 A C ATOM 658 CD2 LEU 115 49.484 47.911 10.530 1.00 30.77 A C ATOM 659 C LEU 115 49.484 47.911 10.530 1.00 30.77 A C ATOM 659 C LEU 115 49.484 47.911 10.530 1.00 30.77 A C ATOM 668 CD2 LEU 116 49.484 47.911 10.530 1.00 30.77 A C ATOM 669 C LEU 116 49.484 47.911 10.530 1.00 30.77 A C ATOM 669 C LEU 116 49.484 47.181 10.043 1.00 31.72 A C ATOM 669 C LEU 116 49.484 47.181 10.043 1.00 31.72 A C ATOM 669 C LEU 116 49.484 47.181 10.043 1.00 30.77 A C ATOM 669 C LEU 116 49.484 47.181 10.043 1.00 30.77 A C ATOM 669 C LEU 116 49.484 47.181 10.043 1.00 30.77 A C ATOM 669 C LEU 116 49.484 47.181 10.043 1.00 30.77 A C ATOM 669 C LEU 116 49.484 47.181 10.043 1.00 30.77 A C ATOM 669 C LEU 116 49.484 47.183 10.048 1.00 28.78 A N A C ATOM 669 C LEU 116 49.486 47.481 1.00 30.77 A C A								
ATOM 641 CE2 PHE 113								
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ATOM 678 N TYR 118 55. 247 46. 663 13. 313 1. 00 27. 75 A N ATOM 679 CA TYR 118 56. 327 45. 796 13. 765 1. 00 29. 68 A C ATOM 680 CB TYR 118 56. 473 44. 586 12. 837 1. 00 29. 52 A C ATOM 681 CG TYR 118 56. 819 44. 903 11. 402 1. 00 28. 58 A C ATOM 682 CD1 TYR 118 55. 922 45. 572 10. 573 1. 00 29. 31 A C ATOM 683 CE1 TYR 118 56. 236 45. 838 9. 239 1. 00 28. 13 A C ATOM 684 CD2 TYR 118 58. 040 44. 510 10. 864 1. 00 28. 81 A C								
ATOM 679 CA TYR 118 56.327 45.796 13.765 1.00 29.68 A C ATOM 680 CB TYR 118 56.473 44.586 12.837 1.00 29.52 A C ATOM 681 CG TYR 118 56.819 44.903 11.402 1.00 28.58 A C ATOM 682 CD1 TYR 118 55.922 45.572 10.573 1.00 29.31 A C ATOM 683 CE1 TYR 118 56.236 45.838 9.239 1.00 28.13 A C ATOM 684 CD2 TYR 118 58.040 44.510 10.864 1.00 28.81 A C								
ATOM 680 CB TYR 118 56.473 44.586 12.837 1.00 29.52 A C ATOM 681 CG TYR 118 56.819 44.903 11.402 1.00 28.58 A C ATOM 682 CD1 TYR 118 55.922 45.572 10.573 1.00 29.31 A C ATOM 683 CE1 TYR 118 56.236 45.838 9.239 1.00 28.13 A C ATOM 684 CD2 TYR 118 58.040 44.510 10.864 1.00 28.81 A C						1.00 29.68	Α	C
ATOM 681 CG TYR 118 56.819 44.903 11.402 1.00 28.58 A C ATOM 682 CD1 TYR 118 55.922 45.572 10.573 1.00 29.31 A C ATOM 683 CE1 TYR 118 56.236 45.838 9.239 1.00 28.13 A C ATOM 684 CD2 TYR 118 58.040 44.510 10.864 1.00 28.81 A C					12.837	1.0029.52		C
ATOM 682 CD1 TYR 118 55.922 45.572 10.573 1.00 29.31 A C ATOM 683 CE1 TYR 118 56.236 45.838 9.239 1.00 28.13 A C ATOM 684 CD2 TYR 118 58.040 44.510 10.864 1.00 28.81 A C			118			1.00 28.58		Č
ATOM 683 CE1 TYR 118 56.236 45.838 9.239 1.00 28.13 A C ATOM 684 CD2 TYR 118 58.040 44.510 10.864 1.00 28.81 A C		682 CD1 TYR						
1110111 001 011 011 011 011 011 011 011	ATOM							
ATOM 685 CE2 TYR 118 58.362 44.769 9.541 1.00 27.91 A C								
	ATOM	685 CEZ TYR	118	58. 362 44. 769	9. 541	1.00 41.91	A	C .

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				FIG. 4-15		(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	686 687 688 689 690 691 692 693 694 695	CZ TYOH TYON ASCA ASCG ASOD1 ASND2 ASC	TR 118 TR 118 TR 118 TR 118 STR 119	57. 459 45. 431 8. 735 1. 00 28. 04 57. 792 45. 681 7. 427 1. 00 29. 86 57. 641 46. 572 13. 863 1. 00 31. 53 57. 683 47. 763 13. 550 1. 00 32. 24 58. 708 45. 903 14. 295 1. 00 32. 40 60. 008 46. 557 14. 459 1. 00 33. 64 60. 511 47. 128 13. 131 1. 00 35. 42 61. 069 46. 066 12. 207 1. 00 36. 36 61. 958 45. 306 12. 584 1. 00 37. 66 60. 560 46. 021 10. 983 1. 00 37. 41 59. 875 47. 697 15. 464 1. 00 34. 07 60. 548 48. 719 15. 348 1. 00 34. 50	A A A A A A A	C O C O N C C C O N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	698 699 700 701 702 703 704 705 706 707 708	CG TY CD1 TY CE1 TY CD2 TY CE2 TY CZ TY OH TY C TY O TY	TR 120	58. 996 47. 514 16. 443 1. 00 33. 92 58. 741 48. 517 17. 472 1. 00 33. 38 57. 510 48. 097 18. 290 1. 00 33. 40 57. 290 48. 870 19. 569 1. 00 33. 30 58. 029 48. 582 20. 715 1. 00 33. 37 57. 818 49. 284 21. 902 1. 00 34. 88 56. 333 49. 886 19. 636 1. 00 33. 62 56. 114 50. 596 20. 813 1. 00 32. 73 56. 859 50. 289 21. 944 1. 00 35. 24 56. 643 50. 977 23. 121 1. 00 37. 51 59. 933 48. 772 18. 396 1. 00 33. 80 60. 472 47. 849 19. 007 1. 00 33. 80 60. 230 18. 401 1. 00 31. 60	A A A A A A A A	N C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	717 718 719 720 721	CG1 VACCG2 VACCG2 VACCG3 VACCG	AL 121 AL 121 AL 121 AL 121 AL 121 YS 122 YS 122 YS 122 YS 122 YS 122	60. 330 50. 038 18. 491 1. 00 31. 69 61. 441 50. 446 19. 343 1. 00 30. 32 62. 672 50. 845 18. 504 1. 00 30. 75 63. 853 51. 140 19. 420 1. 00 28. 68 63. 013 49. 736 17. 525 1. 00 29. 00 61. 008 51. 645 20. 190 1. 00 29. 83 60. 788 52. 738 19. 670 1. 00 30. 47 60. 889 51. 434 21. 495 1. 00 28. 18 60. 464 52. 488 22. 404 1. 00 27. 02 60. 214 51. 910 23. 799 1. 00 23. 73 59. 793 52. 954 24. 819 1. 00 20. 47	A A A A A A A A A A	C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	722 723 724 725 726 727 728 729 730 731 732 733 734	NZ L'C		59. 078 53. 406 27. 174 1. 00 19. 23 60. 062 54. 510 27. 346 1. 00 18. 20 61. 460 53. 635 22. 528 1. 00 27. 64 62. 658 53. 464 22. 315 1. 00 28. 10 60. 947 54. 813 22. 860 1. 00 27. 23 61. 791 55. 979 23. 071 1. 00 27. 82 61. 607 57. 034 21. 974 1. 00 28. 29 62. 537 58. 227 22. 164 1. 00 28. 94 62. 339 59. 308 21. 131 1. 00 29. 91 61. 218 59. 744 20. 889 1. 00 30. 94 61. 385 56. 545 24. 428 1. 00 26. 89 61. 837 56. 036 25. 453 1. 00 27. 03	A A A A A A A A A A A	C N C O N C C C O N C

(Continued)

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ATOM	735	N	TRP	124	60.522	57. 564	24. 444	1.00 23.89	· A	N.
ATOM	736	CA	TRP	124	60.081	58.149	25.713	1.00 24.21	Α	C
ATOM	737	CB	TRP	124	59. 886	59.665	25. 572	1.00 23.25	Α	C
					61.052	60.357	24. 934	1.00 19.79	Ä	Č
ATOM	738	CG	TRP	124						č
ATOM	739		TRP	124	62.444	60.061	25. 127	1.00 19.03	A	
ATOM	740	CE2	TRP	124	63. 175	60.913	24. 270	1.00 19.13	A	C
ATOM	741	CE3	TRP	124	63. 143	59. 157	25.936	1.00 15.51	Α	C
ATOM	742		TRP	124	60.999	61.350	24.006	1.00 18.84	Α	С
ATOM	743	NE1	TRP	124	62. 270	61.690	23.597	1.00 18.74	Α	N
			TRP	124	64. 571	60. 885	24. 196	1.00 17.77	Ā	Ċ
ATOM	744						25. 860	1.00 15.41	Ä	Č
ATOM	745		TRP	124	64. 533	59. 129		1.00 13.41	A	č
ATOM	746	CH2	TRP	124	65. 229	59. 986	24.996			
ATOM	747	C	TRP	124	58. 787	57. 494	26. 209	1.00 24.57	A	C
ATOM	748	0	TRP	124	58. 490	56.350	25.861	1.00 25.71	A	0
ATOM	749	N	ARG	125	58.013	58. 218	27.013	1.00 24.36	Α	N
ATOM	750	CA	ARG	125	56.779	57.670	27.567	1.00 23.36	Α	C
ATOM	751	CB	ARG	125	56. 189	58.621	28.609	1.00 23.81	Α	С
ATOM	752	CG	ARG	125	54. 953	58.065	29.308	1.00 23.85	Α	C
ATOM	753	CD	ARG	125	54. 273	59. 129	30. 143	1.00 26.24	A	Č
			ARG	125	55.090	59. 579	31. 269	1.00 25.99	A	Ň
ATOM	754	NE			55. 293	58. 867	32. 372	1.00 26.04	A	Č
ATOM	755	CZ	ARG	125						N
ATOM	756	NH1	ARG	125	56.051	59. 357	33. 347	1.00 24.42	A	
ATOM	757	NH2	ARG	125	54. 735	57.668	32.500	1.00 25.19	A	N
ATOM	758	C	ARG	125	55. 706	57. 324	26. 541	1.00 24.00	A	C
ATOM	759	0	ARG	125	54.935	56. 387	26.752	1.00 25.04	Α	0
ATOM	760	N	HIS	126	55.651	58.063	25.436	1.00 23.33	Α	N
ATOM	761	CA	HIS	126	54.649	57.800	24.403	1.00 22.86	Α	С
ATOM	762	CB	HIS	126	53.649	58.943	24.353	1.00 21.14	Α	C
ATOM	763	CG	HIS	126	52.987	59. 224	25.662	1.00 22.35	A	C
ATOM	764		HIS	126	53.027	60.316	26. 463	1.00 21.51	A	Č
				126	52. 137	58. 329	26. 274	1.00 22.03	Ä	N
ATOM	765		HIS							C
ATOM	766		HIS	126	51.679	58. 859	27. 395	1.00 23.59	A	
ATOM	767		HIS	126	52. 202	60.064	27. 532	1.00 22.48	A	N
ATOM	768	C	HIS	126	55. 222	57. 599	22.995	1.00 24.43	A	C
ATOM	769	0	HIS	126	54. 599	56.947	22. 153	1.00 23.99	Α	0
ATOM	770	N	SER	127	56.401	58. 163	22.744	1.00 23.89	Α	N
ATOM	771	CA	SER	127	57.039	58.072	21.434	1.00 24.38	Α	C
ATOM	772	CB	SER	127	58.050	59. 213	21.267	1.00 23.49	Α	C
ATOM	773	0G	SER	127	58.909	59.311	22.387	1.00 23.05	Α	0
ATOM	774	C	SER	127	57.737	56. 748	21.146	1.00 24.40	Ā	Č
			SER	127	58. 167	56.050	22.061	1.00 26.55	Ä	ŏ
ATOM	775	0	TYR					1.00 20.00	A	N
ATOM	776	N		128	57.841	56. 420	19.861			
ATOM	777	CA	TYR	128	58. 501	55. 207	19.403	1.00 22.06	A	C
ATOM	778	CB	TYR	128	57. 787	53. 962	19. 928	1.00 21.99	A	C
ATOM	779	CG	TYR	128	56.413	53.712	19. 331	1.00 22.49	A	C
ATOM	780	CD1	TYR	128	55.257	54.112	20.003	1.00 23.20	A	C
ATOM	781		TYR	128	53.992	53.857	19.487	1.00 19.81	Α	C.
ATOM	782		TYR	128	56.267	53.049	18.109	1.00 20.70	Α	C
ATOM	783	CE2	TYR	128	55.007	52. 791	17. 580	1.00 20.87	Α	C
1110111	, 00	720			531001	52. 101	000			

			FIG. 4-17	(Continued)
ATOM	784 CZ TYR	128	53. 872 53. 197 18. 279 1. 00 22. 39	A C
ATOM	785 OH TYR	128	52.614 52.946 17.776 1.00 19.88	A 0
ATOM	786 C TYR	128	58. 509 55. 160 17. 882 1. 00 22. 84	A C
ATOM	787 0 TYR	128	57. 800 55. 922 17. 224 1. 00 24. 63	A 0
ATOM	788 N THR	129	59. 328 54. 281 17. 320 1. 00 22. 88	A N
ATOM	789 CA THR	129	59. 360 54. 125 15. 874 1. 00 25. 24	A C
ATOM	790 CB THR	129	60. 723 54. 474 15. 245 1. 00 27. 54	A C
ATOM	791 OG1 THR	129	61.756 53.676 15.844 1.00 33.01	A 0
ATOM	792 CG2 THR	129	61.025 55.951 15.419 1.00 28.79	A C
ATOM	793 C THR	129	59. 062 52. 675 15. 580 1. 00 24. 85	A C
ATOM	794 0 THR	129	59. 168 51. 811 16. 457 1. 00 22. 29	A 0
ATOM	795 N ALA	130	58.692 52.411 14.337 1.00 24.54	A N
ATOM	796 CA ALA	130	58. 356 51. 062 13. 943 1. 00 25. 98	A C
ATOM	797 CB ALA	130	57. 061 50. 636 14. 618 1. 00 22. 73	A C
ATOM	798 C ALA	130	58. 195 50. 983 12. 445 1. 00 26. 81	A C
ATOM	799 O ALA	130	58. 277 51. 988 11. 740 1. 00 27. 92	A 0
ATOM	800 N SER	131	57. 978 49. 767 11. 965 1. 00 27. 15	A N
ATOM	801 CA SER	131	57. 759 49. 540 10. 556 1. 00 27. 62	A C
ATOM	802 CB SER	131	58. 643 48. 403 10. 059 1. 00 28. 58	A C
ATOM	803 OG SER	131	59. 995 48. 822 10. 022 1. 00 29. 90	A 0
ATOM	804 C SER	131	56. 290 49. 187 10. 426 1. 00 27. 17	A C
ATOM	805 O SER	131	55. 651 48. 779 11. 397 1. 00 27. 00	A O
ATOM	806 N TYR	132	55. 747 49. 351 9. 232 1. 00 27. 56	A N
ATOM	807 CA TYR	132	54. 341 49. 061 9. 029 1. 00 28. 28	A C
ATOM	808 CB TYR	132	53. 532 50. 357 9. 156 1. 00 27. 16	A C
ATOM	809 CG TYR	132	53.649 51.046 10.507 1.00 25.23	A C
ATOM	810 CD1 TYR	132	52.692 50.842 11.500 1.00 24.00	A C
ATOM	811 CE1 TYR	132	52. 790 51. 483 12. 735 1. 00 23. 00	A C
ATOM ATOM	812 CD2 TYR 813 CE2 TYR	132	54. 714 51. 908 10. 785 1. 00 22. 89 54. 822 52. 549 12. 016 1. 00 21. 43	A C
ATOM	814 CZ TYR	132 132	54. 822 52. 549 12. 016 1. 00 21. 43 53. 856 52. 333 12. 985 1. 00 22. 58	A C A C
ATOM	815 OH TYR	132	53. 940 52. 976 14. 198 1. 00 21. 69	
ATOM	816 C TYR	132	54.071 48.418 7.680 1.00 28.72	A O A C
ATOM	817 O TYR	132	54.794 48.639 6.712 1.00 29.54	A O
ATOM	818 N ASP	133	53.028 47.604 7.631 1.00 29.99	A N
ATOM	819 CA ASP	133	52.629 46.956 6.392 1.00 31.05	A C
ATOM	820 CB ASP	133	53.147 45.519 6.314 1.00 31.90	A C
ATOM	821 CG ASP	133	54.541 45.436 5.721 1.00 33.92	A Č
ATOM	822 OD1 ASP	133	54.773 46.042 4.649 1.00 33.52	A O
ATOM	823 OD2 ASP	133	55.400 44.756 6.321 1.00 35.83	A O
ATOM	824 C ASP	133	51. 125 46. 952 6. 334 1. 00 30. 39	A C
ATOM	825 O ASP	133		. A 0
ATOM	826 N ILE	134	50.579 47.598 5.315 1.00 28.05	A N
ATOM	827 CA ILE	134	49. 144 47. 652 5. 157 1. 00 25. 68	A C
ATOM	828 CB ILE	134	48. 732 48. 816 4. 269 1. 00 23. 81	Ä Č
ATOM	829 CG2 ILE	134	47. 221 48. 954 4. 289 1. 00 22. 12	A C
ATOM	830 CG1 ILE	134	49. 421 50. 095 4. 752 1. 00 23. 64	A C
ATOM	831 CD1 ILE	134	49. 232 51. 277 3. 846 1. 00 22. 40	A C
ATOM	832 C ILE	134	48. 635 46. 368 4. 524 1. 00 27. 46	A C

			~		E I (~ 1 -	. 1 Q			(Continue	ed)
			•		r I (G. 4-	. 10				
ATOM ATOM	833 834		ILE TYR	134 135	49. 171 47. 599	45.894 45.805	3. 521 5. 127	1.00 27.19 1.00 29.43	A A	0 N	
ATOM ATOM	835 836		TYR TYR	135 135	46. 985 46. 800	44. 588 43. 588	4. 628 5. 772	1.00 30.54 1.00 33.25	A A	C C	
ATOM ATOM	837 838	CG CD1	TYR TYR	135 135	46. 276 47. 113	42. 242 41. 311	5. 343 4. 731	1.00 35.66 1.00 37.89	A A	C C	
ATOM	839	CE1	TYR	135	46.634	40.068	4.319	1.00 40.13	A	C	
ATOM ATOM	840 841	CD2 CE2		135 135	44. 939 44. 444	41.903 40.666	5. 535 5. 126	1.00 37.34 1.00 40.17	A A	C C	•
ATOM	842	CZ	TYR	135	45.296	39. 751	4.518	1.00 41.67	A	C	
ATOM ATOM	843 844	OH C	TYR TYR	135 - 135	44. 811 45. 629	38. 526 44. 990	4. 105 4. 057	1.00 42.54 1.00 30.05	A A	C 0	
ATOM	845	0	TYR	135	44. 870 45. 341	45. 705 44. 536	4. 704 2. 841	1.00 28.31 1.00 31.33	A A	0 N	
ATOM ATOM	846 847	N CA	ASP ASP	136 136	44.083	44.837	2.168	1.00 33.02	Α	C	
ATOM	848 849	CB CG	ASP ASP	136 136	44. 323 43. 057	44. 857 45. 095	0. 655 -0. 146	1.00 32.51 1.00 33.01	A A	C C	
ATOM ATOM	850	0D1	ASP	136	43.115	45.872	-1.121	1.00 31.21	Α	0	
ATOM ATOM	851 852	OD2 C	ASP ASP	136 ··· 136	42.009 43.019	44. 500 43. 797	0.181 2.549	1.00 34.97 1.00 35.55	A A	0 C	
ATOM	853	0	ASP	136	42.822	42.810	1.846	1.00 36.12	A	0	
ATOM ATOM	854 855	N CA	LEU LEU	137 137	42. 341 41. 303	44. 040 43. 150	3.669 4.192	1.00 38.03 1.00 40.58	A A	N C	
ATOM	856	CB	LEU	137	40.445	43.892	5. 225 6. 477	1.00 40.10 1.00 39.13	A A	C C	
ATOM ATOM	857 858	CG CD1	LEU LEU	137 137	41.160 40.206	44. 413 45. 257	7.307	1.00 37.54	A	C	
MOTA	859 860	CD2	LEU LEU	137 137	41.686 40.392	43. 243 42. 536	7. 286 3. 134	1.00 38.91 1.00 42.88	A A	C C	
ATOM ATOM	861	0	LEU	137	40.038	41.362	3. 225	1.00 43.41	A	0	
ATOM ATOM	862 863	N CA	ASN ASN	138 138	39. 997 39. 132	43. 322 42. 796	2. 141 1. 093	1.00 45.42 1.00 48.50	A A	N C	
ATOM	864	CB	ASN	138	38. 537	43.936	0.264	1.00 49.71	Α	C	
ATOM ATOM	865 866	CG OD1	ASN ASN	138 138 ·	37. 127 36. 873	44. 291 44. 555	0.697 1.871	1.00 50.83 1.00 51.97	A A	C 0	
ATOM	867	ND2	ASN	138	36. 202	44. 296	-0.254	1.00 52.74	A	N	
MOTA MOTA	868 869	C 0	ASN ASN	138 138	39. 884 39. 642	41.824 40.619	0. 191 0. 240	1.00 49.47 1.00 50.62	A A	C 0	
ATOM	870	N	LYS	139	40.794	42.346	-0.626	1.00 50.26	Α	N	
ATOM ATOM	871 872	CA CB	LYS LYS	139 139	41.581 42.510	41.507 42.374	-1.526 -2.382	1.00 51.09 1.00 51.15	A A	C C	
ATOM	873	CG	LYS	139	41.785	43.427	-3. 212 -2. 074	1.00 53.38 1.00 54.25	A A	C C	
ATOM ATOM	874 875	CD CE	LYS LYS	139 139	42. 753 43. 550	44. 331 43. 564	−3. 974 −5. 021	1.00 56.31	A	С	
ATOM	876	NZ C	LYS LYS	139 139	44. 447 42. 413	44. 453 40. 528	-5. 817 -0. 703	1.00 56.39 1.00 51.63	A A	N C	
ATOM ATOM	877 878	0	LYS	139	43.148	39.708	-1.251	1.00 51.80	Α	0	
ATOM ATOM	879 880		ARG	140 · 140	42. 288 43. 025		0. 618 1. 534	1.00 51.49 1.00 51.71	A A	N C	
ATOM	881	CB	ARG	140		38. 408	1. 642	1.00 53.88	A	č	

		•	FIG. 4-19	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	882 CG ARG 883 CD ARG 884 NE ARG 885 CZ ARG 886 NH1 ARG 887 NH2 ARG 888 C ARG 889 O ARG 890 N GLN 891 CA GLN 892 CB GLN 893 CG GLN 894 CD GLN	140 140 140 140 140 140 140 141 141 141	40. 911 38. 495 2. 157 1. 00 57. 36 40. 257 37. 128 2. 211 1. 00 60. 02 40. 936 36. 235 3. 142 1. 00 62. 76 40. 633 34. 950 3. 294 1. 00 64. 87 39. 661 34. 409 2. 570 1. 00 66. 83 41. 298 34. 206 4. 169 1. 00 65. 62 44. 464 39. 603 1. 066 1. 00 50. 29 44. 992 38. 496 1. 002 1. 00 50. 21 45. 096 40. 723 0. 741 1. 00 49. 82 46. 473 40. 707 0. 268 1. 00 48. 70 46. 487 40. 815 -1. 260 1. 00 50. 32 47. 774 40. 348 -1. 909 1. 00 55. 02 47. 640 40. 179 -3. 413 1. 00 57. 33	A C A C A N A C A O A N A C A C A C A C A C A C A C A C A C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	895 OE1 GLN 896 NE2 GLN 897 C GLN 898 O GLN 899 N LEU 900 CA LEU 901 CB LEU 902 CG LEU 903 CD1 LEU 904 CD2 LEU 905 C LEU	141 141 141 142 142 142 142 142 142 142	48. 582 39. 756 -4. 088 1. 00 57. 97 46. 465 40. 509 -3. 947 1. 00 58. 85 47. 293 41. 837 0. 898 1. 00 46. 02 46. 761 42. 880 1. 274 1. 00 45. 33 48. 594 41. 610 1. 013 1. 00 43. 34 49. 505 42. 578 1. 605 1. 00 41. 50 50. 638 41. 824 2. 296 1. 00 41. 17 51. 489 42. 501 3. 359 1. 00 42. 33 52. 443 41. 463 3. 922 1. 00 42. 24 52. 254 43. 677 2. 772 1. 00 42. 66 50. 062 43. 498 0. 520 1. 00 40. 87	A O N A C A C A C A C A C A C A C A C A C A
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	906 O LEU 907 N ILE 908 CA ILE 909 CB ILE 910 CG2 ILE 911 CG1 ILE 912 CD1 ILE 913 C ILE 914 O ILE 915 N THR 916 CA THR 917 CB THR	142 143 143 143 143 143 143 143 144 144	50. 557 43. 030 -0. 506 1. 00 41. 57 49. 978 44. 806 0. 748 1. 00 39. 20 50. 466 45. 789 -0. 217 1. 00 37. 17 49. 921 47. 202 0. 104 1. 00 36. 58 50. 486 48. 225 -0. 874 1. 00 35. 56 48. 398 47. 197 0. 030 1. 00 34. 64 47. 777 48. 494 0. 468 1. 00 37. 28 51. 985 45. 843 -0. 209 1. 00 36. 63 52. 603 45. 859 0. 849 1. 00 36. 63 52. 592 45. 882 -1. 386 1. 00 35. 40 54. 046 45. 933 -1. 459 1. 00 35. 79 54. 616 44. 654 -2. 124 1. 00 35. 59	A
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	918 OG1 THR 919 CG2 THR 920 C THR 921 O THR 922 N GLU 923 CA GLU 924 CB GLU 925 CG GLU 926 CD GLU 927 OE1 GLU 928 OE2 GLU 929 C GLU 930 O GLU	144 144 144 145 145 145 145 145 145 145	54. 192 44. 592 -3. 491 1. 00 37. 13 54. 121 43. 415 -1. 403 1. 00 33. 21 54. 515 47. 152 -2. 243 1. 00 35. 43 55. 700 47. 311 -2. 511 1. 00 36. 45 53. 577 48. 015 -2. 602 1. 00 36. 27 53. 891 49. 214 -3. 369 1. 00 36. 32 52. 962 49. 297 -4. 586 1. 00 38. 36 53. 553 48. 748 -5. 875 1. 00 42. 66 54. 667 49. 639 -6. 418 1. 00 45. 91 55. 745 49. 705 -5. 779 1. 00 45. 49 54. 456 50. 283 -7. 476 1. 00 45. 56 53. 775 50. 496 -2. 544 1. 00 35. 06 52. 874 50. 635 -1. 715 1. 00 34. 22	A

			***					(Continued)
			FIC	G. 4-	20	•.		
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	931 N GLU 932 CA GLU 933 CB GLU 934 CG GLU 935 CD GLU 936 OE1 GLU 937 OE2 GLU 938 C GLU 939 O GLU 940 N ARG 941 CA ARG 942 CB ARG 942 CB ARG 944 CD ARG 945 NE ARG 946 CZ ARG 947 NH1 ARG 948 NH2 ARG 948 NH2 ARG 949 C ARG 949 C ARG 941 N ILE 950 O ARG 951 N ILE 952 CA ILE 953 CB ILE 953 CB ILE 954 CG2 ILE 955 CG1 ILE 955 CG1 ILE 957 C ILE 957 C ILE 958 O ILE 957 C ILE 958 O ILE 957 C ILE 958 O ILE 959 N PRO 960 CD PRO 961 CA PRO 962 CB PRO 963 CG PRO 964 C PRO 965 O PRO 966 N ASN 967 CA ASN	146 146 146 146 146 146 147 147 147 147 147 147 148 148 148 148 148 149 149 149 149 149 150 150	F I 6 54. 692 54. 699 53. 594 53. 708 54. 495 55. 309 54. 495 55. 185 55. 992 55. 376 55. 999 57. 415 58. 271 57. 844 59. 546 55. 066 55. 430 54. 537 55. 096 55. 116 55. 096 55. 430 54. 537 55. 096 55. 440 55. 623 56. 440 55. 623 56. 440 55. 623 56. 440 55. 623 56. 440 55. 623 56. 540 55. 623 56. 55. 096 55. 116 55. 107 56. 879 57. 240 57. 735 57. 443 59. 765 58. 621 58. 621 58. 621 58. 621	51. 428 52. 706 53. 608 53. 924 54. 651 55. 129 54. 754 52. 521 53. 172 51. 638 51. 357 50. 107 48. 821 47. 649 47. 539 46. 143 46. 737 52. 483 53. 364 54. 201 53. 642 53. 642 53. 642 53. 656 54. 557 55. 244 56. 244 57. 56. 244 57. 56. 244 57. 56. 244 57. 56. 244 57. 56. 244 57. 665 57. 665 57. 665 57. 665 57. 665 57. 665 57. 665 57. 666 57. 666	2 0 -2. 782 -2. 079 -2. 630 -4. 107 -4. 455 -3. 528 -5. 660 -0. 579 0. 031 1. 437 1. 774 1. 262 1. 963 1. 650 2. 356 3. 421 1. 987 2. 363 2. 002 3. 568 4. 555 5. 798 6. 940 5. 427 6. 533 4. 891 5. 068 4. 974 4. 930 5. 282 5. 151 5. 614 6. 642 7. 567 6. 748 8. 016	1.00 33.82 1.00 32.54 1.00 33.18 1.00 33.14 1.00 35.19 1.00 35.19 1.00 32.26 1.00 32.38 1.00 30.84 1.00 39.94 1.00 31.91 1.00 34.66 1.00 37.64 1.00 39.76 1.00 40.68 1.00 39.76 1.00 29.74 1.00 26.79 1.00 25.21 1.00 25.21 1.00 24.62 1.00 25.09 1.00 23.14 1.00 23.82 1.00 24.99 1.00 25.52 1.00 24.99 1.00 24.99 1.00 25.52 1.00 24.99 1.00 24.99 1.00 25.52 1.00 24.99 1.00 25.52 1.00 24.99 1.00 25.52 1.00 24.99 1.00 25.52 1.00 24.99 1.00 25.52 1.00 25.52 1.00 25.52 1.00 26.52 1.00 27.47 1.00 29.59 1.00 27.47 1.00 29.59 1.00 29.59 1.00 30.82	A A A A A A A A A A A A A A A A A A A	(Continued) N C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM	967 CA ASN 968 CB ASN 969 CG ASN	150 150 150	60. 950 62. 154 61. 775	52. 064 51. 131 49. 805	8. 016 7. 830 7. 189	1.00 30.82 1.00 32.43 1.00 35.16	A A A	
ATOM ATOM ATOM ATOM ATOM	970 OD1 ASN 971 ND2 ASN 972 C ASN 973 O ASN 974 N ASN	150 150 150 150 150	60. 749 62. 612 61. 336 61. 583 61. 387	49. 215 49. 319 53. 245 54. 348 53. 022	7. 530 6. 271 8. 900 8. 394 10. 208	1.00 36.40 1.00 36.52 1.00 30.50 1.00 31.20 1.00 28.46	A A A A	N C O N
ATOM ATOM ATOM ATOM ATOM	975 CA ASN 976 CB ASN 977 CG ASN 978 OD1 ASN 979 ND2 ASN	151 151 151 151 151	61. 734 63. 137 64. 213 64. 360 64. 965	54. 078 54. 622 53. 571 52. 678 53. 666	11. 154 10. 877 11. 048 10. 219 12. 139	1. 00 28. 87 1. 00 30. 74 1. 00 34. 06 1. 00 36. 24 1. 00 37. 62	A A A A	C C C O N

		•								(Cont	inued)
					FΙ	G. 4	- 21			(Cont	iiiueu/
ATOM	980	С	ASN	151	60. 734	55. 230	11.111	1.00 28.16	A	С	
ATOM	981	0	ASN	151	61.118	56. 400	11.112	1.00 28.85	Ä	Ŏ	
ATOM	982	Ň	THR	152	59. 450		11.064	1.00 26.20	A	N	
ATOM	983	CA	THR	152	58. 415	55. 911	11.041	1.00 24.74	A	C	
ATOM	984	CB	THR	152	57. 119	55. 389	10.399	1.00 25.27	A	Č	
ATOM	985	0G1		152	57. 351	55. 125	9.009	1.00 24.18	A	ŏ	
ATOM	986		THR	152	56.004		10.538	1.00 23.99	A	Č	
ATOM	987	C	THR	152	58. 139	56. 319	12.474	1.00 23.46	Ä	č	
ATOM	988	Ŏ	THR	152	57. 933	55. 476	13. 340	1.00 25.16	A	ŏ	
ATOM	989	Ň	GLN	153	58. 134	57. 620	12. 721	1.00 22.30	A	Ň	
ATOM	990	CA	GLN	153	57.916	58. 129	14.063	1.00 20.67	Ä	Ċ	
ATOM	991	CB	GLN	153	58. 501	59. 534	14.161	1.00 19.09	Ä	č	
ATOM	992	CG	GLN	153	60.002		13.906	1.00 13.74	Ä	Č	•
ATOM	993	CD	GLN	153	60.495	60.853	13.331	1.00 14.57	Ä	Č	
ATOM	994	0E1	GLN	153	60.089	61.260	12. 233	1.00 12.70	A	0	
ATOM	995	NE2	GLN	153	61.375	61.524	14.066	1.00 10.81	Α	N	
ATOM	996	C	GLN	153	56.460	58.112	14.495	1.00 20.53	Α	C	
ATOM	997	0	GLN	153	56. 163	57.979	15.683	1.00 19.36	Α	0	
ATOM	998	N	TRP	154	55.556	58. 229	13.531	1.00 20.90	A	N	
ATOM	999	CA	TRP	154	54.131	58. 213	13.831	1.00 21.02	A	C	
ATOM	1000	CB	TRP	154	53. 733	59. 498	14.550	1.00 22.43	Α	С	
ATOM	1001	CG	TRP	154	52. 312	59. 530	14. 923	1.00 21.90	Α	C	
ATOM	1002		TRP	154	51.695	58. 791	15. 976	1.00 22.22	Α	C	
ATOM	1003		TRP	154	50. 315	59.087	15.942	1.00 23.62	Α	C	
ATOM	1004		TRP	154	52.173	57.902	16. 947	1.00 22.95	Α	C	
ATOM	1005		TRP	154	51.321	60. 228	14. 308	1.00 24.44	Α	C	
ATOM	1006		TRP	154	50.112	59.968	14. 912	1.00 24.78	Α	N	
ATOM	1007		TRP	154	49. 404	58. 526	16.842	1.00 22.94	Α	С	
ATOM	1008		TRP	154	51.263	57. 339	17. 847	1.00 22.07	Α	С	
ATOM	1009		TRP	154	49. 897	57.656	17. 784	1.00 23.43	A	С	
ATOM	1010	C	TRP	154	53. 291	58. 054	12.576	1.00 21.43	A	C	
ATOM	1011	0	TRP	154	53. 642	58. 572	11.518	1.00 22.33	A	0	
ATOM	1012	N	VAL	155	52. 173	57. 343	12. 703	1.00 21.97	A	N	
ATOM	1013	CA	VAL	155	51.267	57. 103		1.00 20.81	A	Ç	
ATOM	1014		VAL	155	51.642	55. 797	10.840	1.00 19.96	A	C	
ATOM ATOM	1015	CG1		155	51.835	54. 687	11.842	1.00 21.34	A	C	
	1016		VAL	155	50.562	55.414	9.833	1.00 20.23	A	C	•
ATOM	1017	C	VAL	155	49. 840	57.004	12. 104	1.00 21.39	A	C	
ATOM	1018	0 N	VAL	155	49.601	56. 425	13. 162	1.00 21.74	A	0	
ATOM	1019	N	THR	156	48. 898	57. 576	11.364	1.00 20.70	A	N	
ATOM	1020	CA	THR	156	47.504	57. 557	11. 768	1.00 21.67	A	C	
ATOM ATOM	1021 1022	CB	THR THR	156 156	47.189	58. 736	12.716	1.00 22.79	A	C	
ATOM ATOM	1022	CG2		156 156	45.771	58. 848	12.890	1.00 25.50	` A	0	
ATOM	1023	CG2	THR	156	47.707	60.031	12. 145	1.00 22.46	A	C	
ATOM	1024	0	THR	156	46.558	57. 633	10.577	1.00 22.20	A	C	
ATOM	1025	N	TRP	157	46.861 45.413	58. 276	9.577	1.00 22.72	A	0 N	
ATOM	1027	CA	TRP	157	40.413	56. 966 56. 985	10. 689 9. 627	1.00 21.38 1.00 21.45	A A	N C	
ATOM	1028	CB	TRP	157	43. 426	55. 825	9. 765	1.00 21.45	A A	C	•
II OIII	1000	OD	114	101	70. 40U	00.020	3. 100	1.00 61.00	n	v	

· v .	F I	G. 4 - 22		(Continued)
ATOM 1029 CG TRP ATOM 1030 CD2 TRP ATOM 1031 CE2 TRP ATOM 1032 CE3 TRP ATOM 1033 CD1 TRP ATOM 1034 NE1 TRP ATOM 1035 CZ2 TRP ATOM 1036 CZ3 TRP ATOM 1036 CZ3 TRP ATOM 1037 CH2 TRP ATOM 1038 C TRP ATOM 1039 O TRP ATOM 1040 N SER ATOM 1040 N SER ATOM 1041 CA SER ATOM 1042 CB SER ATOM 1043 OG SER ATOM 1044 C SER ATOM 1045 O SER ATOM 1046 N PRO ATOM 1047 CD PRO ATOM 1048 CA PRO ATOM 1049 CB PRO ATOM 1049 CB PRO ATOM 1049 CB PRO ATOM 1049 CB PRO	157 43. 995 157 44. 315 157 44. 843 157 44. 208 157 44. 328 157 44. 838 157 45. 265 157 45. 265 157 45. 149 157 43. 650 157 43. 750 158 42. 889 158 42. 064 158 41. 667 158 41. 208 158 40. 613 159 40. 056 159 40. 136 159 38. 876 159 38. 270 159 39. 427	54. 450 9. 599 53. 800 8. 364 52. 531 8. 686 54. 168 7. 019 53. 571 10. 592 52. 417 10. 052 51. 626 7. 708 53. 267 6. 046 52. 011 6. 397 58. 276 9. 801 58. 917 10. 843 59. 855 8. 889 60. 362 7. 502 59. 377 9. 678 58. 176 9. 781 60. 301 10. 247 61. 762 10. 114 59. 922 11. 029 61. 264 11. 419 62. 214 11. 353	1. 00 18. 96 1. 00 19. 67 1. 00 17. 93 1. 00 20. 82 1. 00 21. 01 1. 00 19. 12 1. 00 19. 76 1. 00 19. 30 1. 00 23. 03 1. 00 25. 03	A C A C A A
ATOM 1051 C PRO ATOM 1052 O PRO ATOM 1052 O PRO ATOM 1053 N VAL ATOM 1054 CA VAL ATOM 1055 CB VAL ATOM 1056 CG1 VAL ATOM 1057 CG2 VAL ATOM 1058 C VAL ATOM 1058 C VAL ATOM 1060 N GLY ATOM 1061 CA GLY ATOM 1061 CA GLY ATOM 1062 C GLY ATOM 1063 O GLY ATOM 1064 N HIS ATOM 1065 CA HIS ATOM 1066 CB HIS ATOM 1066 CB HIS ATOM 1067 CG HIS ATOM 1068 CD2 HIS ATOM 1069 ND1 HIS ATOM 1069 ND1 HIS ATOM 1070 CE1 HIS ATOM 1071 NE2 HIS ATOM 1072 C HIS ATOM 1073 O HIS ATOM 1074 N LYS ATOM 1075 CA LYS ATOM 1076 CB LYS ATOM 1076 CB LYS ATOM 1076 CB LYS ATOM 1076 CB LYS	159 37. 901 159 37. 191 160 37. 878 160 36. 977 160 35. 066 160 34. 834 160 37. 679 161 37. 268 161 37. 268 161 37. 268 161 39. 121 161 39. 121 161 39. 144 162 40. 164 162 41. 419 162 41. 614 162 40. 039 162 42. 660 162 43. 636 163 42. 609 163 43. 751 163 43. 372 163 42. 528	59. 090 10. 224 58. 248 10. 771 59. 334 8. 919 58. 640 8. 014 59. 545 7. 689 59. 064 6. 449 59. 559 8. 875 58. 218 6. 730 58. 908 6. 245 57. 080 6. 181 56. 579 4. 962 55. 786 5. 286 55. 950 4. 476 55. 239 4. 695 53. 923 3. 920 54. 875 1. 515 53. 402 1. 874 53. 764 0. 606 54. 656 0. 363 56. 053 4. 305 55. 501 3. 794 57. 364 4. 527 58. 221 4. 224 59. 701 4. 273	1. 00 24. 15 1. 00 25. 36 1. 00 27. 14 1. 00 25. 28 1. 00 23. 99 1. 00 24. 54 1. 00 26. 50 1. 00 26. 15 1. 00 24. 51 1. 00 24. 51 1. 00 24. 05 1. 00 22. 93 1. 00 23. 87 1. 00 24. 24 1. 00 25. 01 1. 00 25. 01 1. 00 25. 86 1. 00 26. 04 1. 00 27. 52 1. 00 27. 58 1. 00 27. 77 1. 00 28. 51 1. 00 28. 82 1. 00 24. 47 1. 00 24. 38 1. 00 24. 47 1. 00 23. 45 1. 00 21. 75 1. 00 21. 55	C C C C C C C C C C C C C C C C C C C

						ז יבו	C 1				(Cor	ntinued)
. == 0. 4							G. 4				_	
ATOM ATOM	1078 1079	CD CE	LYS LYS	163 163		42. 281 41. 464	61. 706 62. 316	3. 335 2. 228	1.00 20.23 1.00 18.07	. A	C C	
ATOM	1080	NZ	LYS	163		41. 315	63. 778	2. 422	1.00 20.95	A	N	
ATOM	1081	C	LYS	163		44. 781	57. 961	5. 309	1.00 23.44	A	Ċ	
ATOM	1082	Ö	LYS	163		44. 425	57.600	6. 433	1.00 23.42	A	Ŏ	
ATOM	1083	N	LEU	164		46.053	58.146	4.979	1.00 23.11	Α	N	
ATOM	1084	CA	LEU	.164		47.117	57.937	5.950	1.00 23.65	Α	C	
ATOM	1085	CB	LEU	164		48.014	56.773	5. 524	1.00 24.35	Α	C	
ATOM	1086	CG	LEU	164		47. 551	55. 351	5. 848	1.00 25.57	A	C	
ATOM	1087		LEU	164		48. 519	54. 349	5. 219	1.00 25.59	A	C	
ATOM	1088		LEU	164		47. 497	55. 162	7. 359	1.00 25.62	A	C	•
ATOM	1089	C	LEU	164		47.970	59. 182	6. 120	1.00 23.21	A	C	
ATOM ATOM	1090 1091	O N	LEU ALA	164 165		48. 175 48. 456	59. 943 59. 383	5. 177 7. 335	1.00 24.34 1.00 21.88	A	O N	
ATOM	1091	CA	ALA	165		49. 319	60.508	7. 649	1.00 21.58	A A	C	
ATOM	1093	CB	ALA	165		48. 548	61.583	8. 376	1.00 21.33	A	Č	,
ATOM	1094	C	ALA	165		50.406	59.953	8. 545	1.00 22.07	A	č	
ATOM	1095	Ŏ	ALA	165		50.115	59. 285	9. 537	1.00 22.91	A	Ŏ	
ATOM	1096	N	TYR	166		51.661	60.208	8. 201	1.00 22.02	A	N	
ATOM	1097	CA	TYR	166		52.745	59.697	9.024	1.00 21.73	A	C	
ATOM	1098	CB	TYR	166		53. 185	58.319	8.520	1.00 22.38	Α	С	
ATOM	1099	CG	TYR	166		53.814	58. 315	7. 141	1.00 22.11	Α	C	
ATOM	1100	CD1	TYR	166		55. 148	58.661	6.964	1.00 21.28	A	C	
ATOM	1101	CE1		166		55. 733	58. 638	5. 704	1.00 22.05	A	C	
ATOM	1102		TYR	166		53.074	57. 949	6.015	1.00 20.67	A	C	
ATOM	1103	CE2		166		53.648	57. 923	4. 753	1.00 20.02	A	C	
ATOM ATOM	1104 1105	CZ OH	TYR TYR	166 166		54. 981 55. 566	58. 268 58. 252	4. 603 3. 352	1.00 21.75 1.00 20.77	A	C	
ATOM	1105	C	TYR	166		53. 927	60. 643	9. 057	1.00 20.77	A A	0 C	
ATOM	1107	ŏ	TYR	166		54. 108	61.464	8. 157	1.00 21.04	A	. 0	
ATOM	1108	Ň	VAL	167		54. 722	60. 529	10. 111	1.00 20.28	Ä	N	
ATOM	1109	CA	VAL	167		55. 886	61.371	10. 264	1.00 19.16	Ä	Ċ	
ATOM	1110	CB	VAL	167		55. 924	62.011	11.644	1.00 19.56	Ā	Č	
ATOM	1111	CG1	VAL	167		57.103	62.984	11.731	1.00 18.58	Α	C	
ATOM	1112	CG2		167		54.609	62.713	11.916	1.00 18.36	A	C	
ATOM	1113	C	VAL	167		57. 135	60. 537	10.078	1.00 20.06	A	C	
ATOM	1114	0	VAL	167		57. 287	59. 474	10.679	1.00 21.80	A	0	
ATOM	1115	N	TRP	168		58. 030	61.023	9. 233	1.00 19.65	Ą	N	
ATOM	1116	CA	TRP	168		59. 268	60.320	8. 964	1.00 19.61	A	C	
ATOM	1117	CB	TRP	168		59.164	59.558	7.646	1.00 20.07	· A	C	
ATOM ATOM	1118 1119	CC	TRP TRP	168 168		60.387	58. 772	7. 353 6. 300	1.00 23.12 1.00 21.38	A	C	
ATOM	1120		TRP	168	٠.	61.319 62.353	59. 011 58. 061	6. 436	1.00 21.58	A A	C	
ATOM	1121		TRP	168		61.382	59. 936	5. 256	1.00 21.38	Ä	Ċ	
ATOM	1122	CD1		168		60.873	57. 712	8.066	1.00 22.86	A	Č	
ATOM	1123	NE1		168		62.056	57. 281	7. 521	1.00 21.54	Ä	Ň	
ATOM	1124		TRP	168		63. 445	58. 012	5. 563	1.00 23.71	Ä	Ċ	
ATOM	1125	CZ3	TRP	168		62.468	59. 889	4. 386	1.00 23.21	A		
ATOM	1126	CH2	TRP	168		63.484	58. 934	4. 546	1.00 22.74	A	C	

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					T2 T	C 1	9.4			(Con	tinued)
		•			FI	G. 4	- 24				
MOTA	1107	C	ממיד	160	60 406	61.327	8. 906	1.00 19.17	A	С	
ATOM	1127		TRP	168	60. 406	62.319	8. 187	1.00 19.11	A	Õ	
ATOM	1128		TRP	168	60. 331	61.072	9. 682	1.00 19.01	A	N	
ATOM	1129		ASN	169 169	61. 452 62. 589	61.969	9. 732	1.00 13.20	A	Č	
ATOM	1130		ASN	169	63. 374	61.902	8. 417	1.00 21.00	A	Č	
ATOM	1131		ASN ASN	169	64. 056	60. 565	8. 217	1.00 25.33	A	Č	
ATOM	1132	CG OD1		169	64. 410	60.196		1.00 20.24	A	ŏ	
ATOM	1133	ND2		169	64. 255	59.832	9. 307	1.00 27.22	A	N	
ATOM ATOM	1134 1135		ASN	169	62. 122	63.394		1.00 19.72	A	Ċ	
ATOM	1136	0	ASN	169	62. 582	64.344	9. 378	1.00 19.61	Ä	ŏ	
ATOM	1137	N	ASN	170	61. 182	63. 522	10. 938	1.00 19.01	A	Ň	
ATOM	1138	CA	ASN	170	60. 654	64.817	11. 354	1.00 18.95	Ä	Ċ	
ATOM	1139	CB	ASN	170	61.806	65.679	11. 887	1.00 19.76	A	Č	
ATOM	1140	CG	ASN	170	62. 326	65.193	13. 239	1.00 21.23	A	Č	
ATOM	1141	0D1		170	62. 690	64.025	13. 404	1.00 23.29	A	0	
ATOM	1142	ND2		170	62. 362	66.092	14. 210	1.00 21.16	A	N	
ATOM	1143	C	ASN	170	59.828	65.621	10.341	1.00 18.94	Α	C	
ATOM	1144	Ŏ	ASN	170	59. 594	66.815	10.541	1.00 17.99	Α	0	
ATOM	1145	N	ASP	171	59. 385	64.974	9.264	1.00 18.46	Α	N	
ATOM	1146	CA	ASP	171	58. 566	65.643	8. 254	1.00 18.64	A	C	
ATOM	1147	CB	ASP	171	59. 271	65.696	6.898	1.00 18.52	Α	C	
ATOM	1148	CG	ASP	171	60.353	66.750	6.836	1.00 17.77	A	C	
ATOM	1149	OD1		171	60.126	67.876	7. 307	1.00 17.30	A	0	
ATOM	1150	0D2		171	61.436	66.454	6. 294	1.00 24.17	A	0	
ATOM	1151	C	ASP	171	57. 255	64.888	8.099	1.00 20.36	A	Ç	
ATOM	1152	0	ASP	171	57. 182	63.690	8. 382	1.00 21.44	A	0	
ATOM	1153	N	ILE	172	56. 225	65. 585	7. 632	1.00 19.52	A	N	
ATOM	1154	CA	ILE	172	54. 908	64.983	7.466	1.00 18.52	A	C	
ATOM	1155	CB	ILE,		53. 813	65.966	7.899	1.00 18.99	A	C	
ATOM	1156		ILE	172	52. 443	65. 329	7. 734	1.00 17.69	A	C	
ATOM	1157		ILE	172	54.053	66. 394 67. 538	9. 350 9. 795	1.00 18.78 1.00 18.44	A A	C	
ATOM ATOM	1158	CDI	ILE ILE	172 172	53. 167 54. 609	64. 539	6.044	1.00 18.44	A	C	
ATOM	1159 1160	0	ILE	172	54. 905	65. 246	5.085	1.00 19.61	A	. 0	
ATOM	1161	N	TYR	173	54. 017	63. 358	5. 921	1.00 17.61	A	N	
' ATOM	1162	CA	TYR	173	53. 645	62.808	4.625	1.00 16.59	Ä	Ĉ	
ATOM	1163	, CB	TYR	173	54. 519	61.612	4. 256	1.00 14.94	Ä	č	
ATOM	1164	CG	TYR	173	55. 983	61.921	4. 121	1.00 15.66	Ä	Č	
ATOM	1165		TYR	173	56. 815	61.978	5. 237	1.00 16.67	A	Č	
ATOM	1166		TYR	173	58. 170	62. 271	5. 100	1.00 16.34	A	Ċ	
ATOM	1167		TYR	173	56. 541	62.165	2.870	1.00 15.99	Α	C	
ATOM	1168		TYR	173	57.879	62.460	2.727	1.00 13.89	Α	C	
ATOM	1169	CZ	TYR	173	58. 685	62.512	3.838	1.00 15.53	Α	C	
ATOM	1170	OH	TYR	173	60.004	62.837	3.678	1.00 21.66	Α	0	
ATOM	1171	C	TYR	173	52. 198	62.341	4.679	1.00 17.34	A	C	
ATOM	1172	0	TYR	173	51.683	62.008	5.748	1.00 14.56	A	0	
ATOM	1173	N	VAL	174	51.552	62.306	3.518	1.00 18.18	A	Ñ	
ATOM	1174	CA	VAL	174	50.174	61.865	3. 444	1.00 19.46	A	C	
ATOM	1175	CB	VAL	174	49. 212	63.060	3. 319	1.00 18.88	A	C	

1224

ATOM

CG LEU

180

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(Continued) FIG. 4-25 174 47.775 62.564 3.207 C ATOM 1176 CG1 VAL 1.00 19.37 A 63.969 1.00 20.44 C CG2 VAL 49.359 4.534 ATOM 1177 174 A C 60.928 **ATOM** 1178 C VAL 174 49.948 2.268 1.00 21.57 A 61.129 1.00 22.86 A 0 ATOM 1179 0 VAL 174 50.485 1.185 ATOM 1180 N LYS 175 49.154 59.891 2.500 1.00 23.19 A N 48. 824 49. 275 CA LYS 58.934 1.461 1.00 23.86 C **ATOM** 1181 175 A C LYS 57.516 1.831 1.00 24.28 A **ATOM** 1182 CB 175 C CG LYS 50.759 57.352 1.00 28.82 A ATOM 1183 175 2.113 C **ATOM** 1184 CD LYS 175 51.100 55.895 2.422 1.00 29.18 A LYS CE 175 51.107 55.043 1.163 1.00 29.84 C **ATOM** 1185 A ATOM 1186 NZ LYS 175 52.263 55.409 0.291 1.00 31.80 A N 47.314 58.935 1.00 24.49 C LYS 1.338 Α ATOM 1187 175 **ATOM** LYS 46.615 58.606 2.293 1.00 25.05 0 1188 0 175 A 46.820 59.319 0.166 1.00 24.77 ILE ATOM 1189 N N 176 Α CA 45.394 59.327 -0.1021.00 24.70 C ATOM 1190 ILE 176 A CB ILE 45.095 60.028 1.00 22.88 C **ATOM** 1191 176 -1.437A C **ATOM** 1192 CG2 ILE 176 43.605 60.073 -1.6791.00 21.75 A 1193 CG1 ILE 45.677 61.443 -1.4231.00 21.52 C **ATOM** 176 Α CD1 ILE 45.016 62.379 -0.4241.00 23.58 C **ATOM** 1194 176 A C 44.99557.860 1.00 26.89 **ATOM** C ILE 176 -0.211Α 1195 ILE 43.979 57.428 0.328 1.00 26.38 **ATOM** 1196 0 176 A 0 45.829 -0.906**GLU** 57.097 **ATOM** 1197 N 177 1.00 29.47 Α N ATOM 1198 CA GLU 177 45.597 55.672 -1.1041.00 31.88 A C CB GLU 45.412 55.380 1.00 35.29 C **ATOM** 1199 177 -2.594Α ATOM 1200 CG GLU 177 44.308 56.190 -3.2481.00 38.36 C A CD GLU 42.925 55.776 -2.784C **ATOM** 1201 177 1.00 41.13 A **ATOM** 1202 OE1 GLU 41.951 56.495 -3.1051.00 45.06 177 A 0 **ATOM** 1203 OE2 GLU 177 42.810 54.730 -2.1071.00 40.42 A 0 **ATOM** 1204 C **GLU** 177 46.796 54.895 -0.5691.00 31.55. C A 1205 0 GLU 47.940 55.223 -0.8721.00 31.59 **ATOM** 177 0 A **PRO** 46.544 53.840 0.221 **ATOM** 1206 N 178 1.00 31.40 Α N 1.00 30.50 CD 53.240 \mathbb{C} **ATOM** 1207 PRO 178 45.218 0.438 Α CA **PRO** 178 47.591 53.000 0.814 1.00 29.97 C ATOM 1208 A PR0 46.796 1.509 C **ATOM** 1209 CB 178 51.902 1.00 30.05 Α ATOM 1210 CG PRO 178 45.567 51.805 0.684 1.00 31.07 A C 1.00 29.50 C **ATOM** 1211 PR₀ 178 48.633 52.436 -0.150A **PRO** 49.727 52.062 **ATOM** 1212 0 178 0.269 1.00 31.00 0 A 48.308 52.379 1213 ASN -1.4361.00 28.20 ATOM N 179 Α N ATOM 1214 CA ASN 179 49.251 51.838 -2.4091.00 27.53 A C C 1215 CB ASN 179 48.568 50.805 -3.2991.00 26.23 A **ATOM** CG ASN 51.409 -4.144 1.00 25.74 C **ATOM** 1216 179 47.474 A 1.00 26.59 1217 OD1 ASN 46.494 51.948 -3.6260 **ATOM** 179 A 51.329 1.00 26.72 ND2 ASN -5.452A N **ATOM** 1218 179 47.635 **ATOM** 1219 C ASN 179 49.854 52.916 -3.2851.00 27.48 A C ASN 1220 0 179 50.818 52.670 -4.0041.00 28.42 A 0 ATOM LEU 1.00 26.68 N 49.289 54.115 -3.231A N 1221 180 ATOM C -4.050 1.00 26.11 CA 55, 200 A 1222 LEU 180 49.805 ATOM C 1223 CB LEU 48.658 56.125 -4.4561.00 24.86 A ATOM 180

SUBSTITUTE SHEET (RULE 26)

-5.238

47.574 55.370

1.00 25.87

C

1273

ATOM

N.

THR

186

29/246

(Continued) FIG. 4 - 26 C 56.359 -5.8561.00 23.58 46.604 **ATOM** 1225 CD1 LEU 180 C 1.00 22.86 48. 224 54.503 -6.3281226 CD2 LEU 180 **ATOM** C 50.938 55.996 -3.3911.00 25.78 A 1227 C LEU 180 **ATOM** 0 55.883 -2.1851.00 23.62 A 1228 LEU 180 51.185 **ATOM** 0 1229 56.789 -4.194 1.00 24.96 N 51.669 A PR₀ 181 N **ATOM** C 51.687 56.842 -5.6671.00 23.41 A PR₀ 181 ATOM 1230 CD C 1.00 23.35 A 52.766 57.580 -3.634PR₀ 181 **ATOM** 1231 CA C 1.00 22.16 A PR₀ 181 53.403 58.217 -4.870**ATOM** 1232 CB C 1.00 22.72 1233 CG PR₀ 53.124 57.201 -5.944A 181 **ATOM** 1.00 22.15 C 58.613 -2.667Α 52.216 1234 C PR₀ 181 ATOM 1.00 21.88 0 59.173 -2.880A 51.144 0 PR₀ 181 ATOM 1235 N 1.00 21.65 182 52.954 58.864 -1.601Α **ATOM** 1236 N SER C 52.516 59.829 -0.6201.00 20.50 A 1237 CA SER 182 ATOM 1.00 22.61 C 182 52.999 59.404 0.765 Α **ATOM** 1238 CB SER 59.345 0.806 1.00 23.55 A 0 1239 182 54.408 **ATOM** 0GSER 1.00 19.05 Α C 182 53.034 61.222 -0.947**ATOM** 1240 SER C 0 54.003 61.380 -1.6871.00 17.74 A 182 0 SER **ATOM** 1241 N 1.00 17.87 52.366 62.233 -0.402Α **ATOM** 1242 N TYR 183 C -0.6111.00 15.17 A 1243 CA 183 52.786 63.606 ATOM TYR C -0.832 183 51.595 64.523 1.00 12.09 Α ATOM 1244 CB TYR -1.90550.676 64.028 1.00 12.54 C CG TYR 183 Α ATOM 1245 49.729 1.00 8.93 C 183 63.052 -1.625Α CD1 TYR ATOM 1246 Ċ 62.554 -2.6101.00 11.95 A 48.916 CE1 TYR 183 ATOM 1247 C -3.2141248 183 50.782 64.494 1.00 9.42 A **ATOM** CD2 TYR -4.218C 49.961 63.990 1.00 10.27 ATOM 1249 CE2 TYR 183 Α C -3.903TYR 183 49.032 63.019 1.00 10.59 Α 1250 CZ**ATOM** 1.00 14.71 0 1251 OH TYR 183 48.205 62.494 -4.867A **ATOM** C 1252 183 53.532 64.067 0.617 1.00 15.72 A **ATOM** C **TYR** 53.208 1.00 17.69 0 1253 0 183 63.679 1.740 A **TYR** ATOM 54.540 64.893 0.386 1.00 14.64 A N ATOM 1254 N ARG 184 C 184 55.342 65.436 1.452 1.00 14.10 A **ATOM** 1255 CA ARG 56.786 65.593 0.970 1.00 16.84 C **ATOM** 1256 CB **ARG** 184 A 57.725 1.989 1.00 20.48 C ARG 66.203 ATOM 1257 CG 184 A 59.170 65.912 1.629 1.00 20.61 C ATOM 1258 CD ARG A 184 1.00 20.21 NE 60.095 66.485 2.598 Α **ATOM** 1259 ARG 184 ATOM 61.407 66.288 2.583 1.00 19.46 1260 CZ ARG 184 A 1.650 1.00 17.13 NH1 61.954 65.529 A ATOM 1261 ARG 184 3.506 1.00 20.35 1262 62.170 66.853 A N ATOM NH2 ARG 184 C ATOM 1263 C ARG 184 54.736 66.779 1.820 1.00 14.10 Α 1264 54.569 67.650 0.972 1.00 14.71 0 **ATOM** 0 ARG 184 A 54.390 3.089 1.00 15.27 N **ATOM** 1265 N ILE 185 66.937 Α CA 53.804 68.175 3.572 1.00 14.44 Α C 1266 ILE 185 ATOM 67.884 1.00 16.20 C CB ILE 185 52.786 4.692 Α **ATOM** 1267 CG2 ILE 52.091 69.175 5.115 1.00 14.78 C 1268 185 Α ATOM C 4.202 1269 CG1 ILE 185 51.770 66.842 1.00 15.25 A **ATOM** 67.250 2.947 1.00 12.00 C A **ATOM** 1270 CD1 ILE 185 51.021 C 1271 C ILE 185 54.847 69.172 4.091 1.00 14.33 Α ATOM ILE 54.647 70.377 3.994 1.00 14.95 0 **ATOM** 0 185 1272

SUBSTITUTE SHEET (RULE 26)

68.676

4.646

55.950

1.00 14.38

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					(Continued)
				FIG. 4 - 27	
ATOM ATOM	1274 1275	CA THR CB THR	186 186	56. 995 69. 555 5. 169 1. 00 15. 05 A 57. 051 69. 549 6. 717 1. 00 15. 72 A	
ATOM	1276	OG1 THR	186	57. 308 68. 218 7. 181 1. 00 18. 48 A	
ATOM	1277	CG2 THR	186	55.734 70.060 7.323 1.00 13.92 A	
ATOM	1278	C THR	186	58. 384 69. 190 4. 663 1. 00 17. 06 A	
ATOM	1279	0 THR	186	58. 643 68. 055 4. 262 1. 00 19. 33 A	
ATOM ATOM	1280 1281	N TRP CA TRP	187 187	59. 275 70. 174 4. 696 1. 00 18. 28 A 60. 655 70. 020 4. 253 1. 00 16. 04 A	
ATOM	1282	CB TRP	187	60. 655 70. 020 4. 253 1. 00 16. 04 A 60. 843 70. 734 2. 915 1. 00 13. 96 A	
ATOM	1283	CG TRP	187	60.392 69.949 1.736 1.00 14.75 A	
ATOM	1284	CD2 TRP	187	59. 055 69. 841 1. 234 1. 00 15. 37 A	
ATOM	1285	CE2 TRP	187	59. 093 68. 954 0. 135 1. 00 15. 22 A	
ATOM	1286	CE3 TRP	187	57. 829 70. 405 1. 606 1. 00 12. 92 A	
ATOM ATOM	1287 1288	CD1 TRP NE1 TRP	187 187	61. 165 69. 149 0. 941 1. 00 14. 94 A 60. 392 68. 549 -0. 020 1. 00 15. 60 A	
ATOM	1289	CZ2 TRP	187	60. 392 68. 549 -0. 020 1. 00 15. 60 A 57. 949 68. 616 -0. 597 1. 00 17. 91 A	
ATOM	1290	CZ3 TRP	187	56.692 70.074 0.881 1.00 16.75 A	
ATOM	1291	CH2 TRP	187	56. 758 69. 185 -0. 211 1. 00 17. 84 A	· C
ATOM	1292	C TRP	187	61. 607 70. 620 5. 292 1. 00 15. 71 A	
ATOM ATOM	1293 1294	O TRP N THR	187 188	62. 804 70. 725 5. 053 1. 00 19. 54 A	
ATOM	1295	CA THR	188	61. 077 70. 999 6. 449 1. 00 13. 19 A 61. 892 71. 605 7. 493 1. 00 11. 35 A	
ATOM	1296	CB THR	188	61. 122 72. 737 8. 180 1. 00 11. 04 A	
ATOM	1297	OG1 THR	188	59. 835 72. 253 8. 587 1. 00 9. 11 A	
ATOM	1298	CG2 THR	188	60. 955 73. 920 7. 232 1. 00 7. 35 A	. C
ATOM	1299	C THR	188	62. 384 70. 642 8. 572 1. 00 12. 10 A	
ATOM ATOM	1300 1301	0 THR N GLY	188 189	63. 198 71. 016 9. 415 1. 00 9. 49 A 61. 881 69. 412 8. 552 1. 00 14. 44 A	
ATOM	1302	CA GLY	189	61.881 69.412 8.552 1.00 14.44 A 62.296 68.426 9.538 1.00 16.08 A	
ATOM	1303	C GLY	189	63. 794 68. 421 9. 782 1. 00 15. 86 A	
ATOM	1304	0 GLY	189	64.584 68.685 8.881 1.00 17.65 A	
ATOM	1305	N LYS	190	64.196 68.117 11.004 1.00 17.28 A	
ATOM	1306	CA LYS	190	65. 612 68. 096 11. 346 1. 00 18. 87 A	
ATOM ATOM	1307 1308	CB LYS	190 190	66. 189 69. 512 11. 264 1. 00 20. 03 A 67. 679 69. 588 11. 472 1. 00 22. 58 A	_
ATOM	1309	CD LYS	190	67. 679 69. 588 11. 472 1. 00 22. 58 A 68. 181 70. 997 11. 256 1. 00 27. 62 A	_
ATOM	1310	CE LYS	190	69. 698 71. 060 11. 386 1. 00 31. 27 A	
ATOM	1311	NZ LYS	190	70. 207 72. 451 11. 273 1. 00 35. 57 A	
ATOM	1312	C LYS	190	65. 799 67. 530 12. 747 1. 00 18. 55 A	
ATOM ATOM	1313 1314	O LYS N GLU	190	65. 384 68. 134 13. 737 1. 00 18. 41 A	
ATOM	1314	CA GLU	191 191	66. 426 66. 362 12. 811 1. 00 19. 79 A 66. 674 65. 661 14. 062 1. 00 21. 70 A	
ATOM	1316	CB GLU	191	66.674 65.661 14.062 1.00 21.70 A 67.796 64.653 13.851 1.00 23.41 A	
ATOM	1317	CG GLU	191	67. 894 63. 598 14. 937 1. 00 29. 95 A	
ATOM	1318	CD GLU	191	69.018 62.605 14.689 1.00 30.89 A	C
ATOM	1319	OE1 GLU	191	68. 970 61. 497 15. 262 1. 00 33. 70 A	
ATOM	1320 1321	OE2 GLU C GLU	191 191	69. 952 62. 932 13. 929 1. 00 33. 21 A	
ATOM ATOM	1322	O GLU	191	67.015 66.583 15.236 1.00 21.53 A 67.930 67.397 15.156 1.00 22.21 A	
111 OIII	1000	0 000	101	01.000 01.001 10.100 1.00 44.41 A	U

			•		· (Contin	ned)
		•.	FIG. 4	- 28	(Oontain	ucu
		an 100			A N	
ATOM		SP 192	66. 262 66. 45 66. 470 67. 24		A C	
ATOM		SP 192 SP 192	67. 810 66. 88		A Č	
ATOM ATOM		ASP 192 ASP 192	67. 922 65. 40		A C	
ATOM	1320 CO A		66.891 64.77		A O	
ATOM	1328 OD2 A		69.049 64.86		A 0	
ATOM		SP 192	66.425 68.75		A C	
ATOM		SP 192	66.998 69.48		A 0	
ATOM		LE 193	65.748 69.24	2 16.304 1.00 21.66	A N	
ATOM		ILE 193	65.685 70.68		A C	
ATOM		ILE 193	66.747 71.11		A C	
ATOM	1334 CG2 I		66.570 72.56		A C	
ATOM	1335 CG1 I		68.142 70.88		A C	
ATOM		ILE 193	69. 263 71. 19		A C	
ATOM		ILE 193	64.318 71.17		A C	
ATOM		ILE 193	63. 736 72. 06		A O A N	
ATOM		ILE 194	63.814 70.59 62.506 70.96		A N A C	
ATOM		ILE 194 ILE 194	62.596 71.54		A C	
ATOM ATOM	1341 CB I 1342 CG2 I		61. 209 71. 94		A C	
ATOM	1342 CG2 1		63. 551 72. 75		A Č	
ATOM	1344 CD1 I		63.118 73.93		A C	
ATOM		ILE 194	61.663 69.70		A C	
ATOM		ILE 194	62.066 68.71		A 0	
ATOM		TYR 195	60.511 69.72		A N	
ATOM	1348 CA 7	TYR 195	59.592 68.59		A C	
ATOM		TYR 195	59. 338 68. 07		A C	
ATOM		TYR 195	60. 560 67. 77		A C	
ATOM		TYR 195	61. 427 68. 80		A C	
ATOM	1352 CE1 3		62.485 68.55		A C A C A C	
ATOM	1353 CD2 1		60.799 66.49			
ATOM	1354 CE2 7		61.859 66.23 62.694 67.27		A C A C	
ATOM		TYR 195 TYR 195	63.725 67.04		A O	
ATOM ATOM		TYR 195	58. 242 69. 01		A C	
ATOM		TYR 195	57. 574 69. 90		A O	
ATOM		ASN 196	57. 851 68. 38		A N	
ATOM		ASN 196	56. 578 68. 68		A C	
ATOM		ASN 196	56.772 68.89		A C	
ATOM		ASN 196	57. 591 70. 13			
ATOM	1363 OD1	ASN 196	57.132 71.26			•
ATOM	1364 ND2		58.819 69.92			
ATOM		ASN 196	55.686 67.43			
ATOM		ASN 196	56.050 66.34		A 0	
ATOM		GLY 197	54. 522 67. 61			
ATOM		GLY 197	53.622 66.48			
ATOM		GLY 197 GLY 197	53. 880 65. 65 53. 059 64. 79			
ATOM		GLY 197 ILE 198	55. 023 65. 84			
ATOM	1911 1/	100	00.020 00.0		11 11	

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									â	_
ATOM	1372	CA	ILE	198	55. 378	65.097	16. 298	1.00 16.59	Α	C
ATOM	1373	CB	ILE	198	56.425	63.991	16.011	1.00 18.21	Α	C
ATOM	1374	CG2	ILE	198	55.874	63.013	14.987	1.00 18.51	A	C
ATOM	1375	CG1	ILE	198	57. 724	64.602	15. 494	1.00 17.86	A	Č
								1.00 19.35		Č
ATOM	1376	CD1	ILE	198	58. 798	63. 565	15. 214		A	
ATOM	1377	C	ILE	198	55.946	66.057	17.318	1.00 15.95	A	C
ATOM	1378	0	ILE	198	56.507	67.091	16.966	1.00 17.63	Α	0
ATOM	1379	N	THR	199	55.809	65.700	18.583	1.00 15.42	Α	N
ATOM	1380	CA	THR	199	56. 264	66.547	19.672	1.00 16.68	Α	С
ATOM	1381	CB	THR	199	55. 374	66.316	20. 908	1.00 17.40	A	Č
							21.301	1.00 18.82		Õ
ATOM	1382	0G1	THR	199	55. 462	64. 944			A	
ATOM	1383	CG2	THR	199	53.924	66.619	20. 583	1.00 15.72	A	C
ATOM	1384	C	THR	199	, 57. 716	66.334	20.076	1.00 16.00	Α	C
ATOM	1385	0	THR	199	58.317	65.325	19.734	1.00 16.12	Α	0
ATOM	1386	N	ASP	200	58. 276	67.301	20.801	1.00 16.87	A	N
ATOM	1387	CA .	ASP	200	59.649	67. 193	21. 289	1.00 15.49	A	Ĉ
ATOM	1388	CB	ASP	200	60. 315	68. 576	21.418	1.00 14.82	A	č
ATOM	1389	CG	ASP	200	59. 681	69.446	22.491	1.00 17.16	A	C
ATOM	1390		ASP	200	58. 517	69.190	22.873	1.00 16.41	A	0
ATOM	1391		ASP	200	60.348	70.403	22.945	1.00 15.97	Α	0
ATOM	1392	C	ASP	200	59.496	66.515	22.641	1.00 15.54	A	C
ATOM	1393	0	ASP	200	58.388	66.118	22.999	1.00 17.01	Α	0
ATOM	1394	Ň	TRP	201	60.581	66.381	23. 395	1.00 15.10	A	Ň
ATOM	1395	CA	TRP	201	60. 504	65. 699	24. 672	1.00 13.14	A	Ĉ
ATOM	1396	CB	TRP	201	61.885	65.619	25.326	1.00 14.90	A	C
ATOM	1397	CG	TRP	201	61.905	64.679	26.510	1.00 15.25	A	Č
ATOM	1398	CD2	TRP	201	61.412	64.953	27.828	1.00 13.65	A	C
ATOM	1399	CE2	TRP	201	61.500	63. 753	28.564	1.00 13.52	Α	C
ATOM	1400	CE3	TRP	201	60.902	66.096	28.456	1.00 11.78	Α	C
ATOM	1401	CD1	TRP	201	62.269	63.360	26.507	1.00 13.81	A	C
ATOM	1402	NE1	TRP	201	62.025	62.799	27.733	1.00 13.64	A	N
ATOM	1403	CZ2	TRP	201	61.096	63.661	29.897	1.00 14.03	Ä	Č
ATOM	1404	CZ3	TRP	201	60. 502	66.009	29. 778	1.00 12.04		Č
									A	
ATOM	1405	CH2	TRP	201	60.601	64. 797	30.486	1.00 14.87	A	C
ATOM	1406	C	TRP	201	59. 529	66. 327	25.662	1.00 14.42	. A	C
ATOM	1407	0	TRP	201	58. 635	65.656	26. 175	1.00 13.63	. А	0
ATOM	1408	N	VAL	202	59. 691	67.615	25.931	1.00 15.14	Α	N
ATOM	1409	CA	VAL	202	58.830	68.265	26.911	1.00 14.23	Α	C
ATOM	1410	CB	VAL	202	59.402	69.639	27.330	1.00 12.99	Α	C
ATOM	1411		VAL	202	59.010	70.716	26.322	1.00 11.02	Ä	Č
	1412	CG2		202			28.753			č
ATOM					58. 947	69. 963			A	
ATOM	1413	Ç	VAL	202	57. 365	68. 401	26.518	1.00 15.76	A	C
ATOM	1414	0	VAL	202	56. 497	68. 404	27. 391	1.00 18.74	A	0
ATOM	1415	N	TYR	203	57. 072	68.518	25. 226	1.00 15.58	A	N
ATOM	1416	CA	TYR	203	55.676	68.606	24.805	1.00 14.25	A	C
ATOM	1417	CB	TYR	203	55. 556	69.078	23.354	1.00 14.63	Α·	C
ATOM	1418	CG	TYR	203	55. 227	70. 542	23. 227	1.00 12.35	Α	C
ATOM	1419	CD1	TYR	203	56. 231	71.508	23. 193	1.00 11.91	Ä	č
	1420		TYR	203	55. 920	72.867	23. 108	1.00 11.31	Ä	č
ATOM	1460	OLI	111/	200	00. 940	14.00(20. 100	1.00 11.40	n	U

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(Continued)

FIG. 4-30

ATOM	1421	CD2	TYR	203	53.902	70.966	23.177	1.00 12.17	Α	С
ATOM	1422		TYR	203	53. 579	72. 314	23. 099	1.00 10.57	A	C
ATOM	1423	CZ	TYR	203	54. 588	73. 259	23. 061	1.00 9.67	Ä.	Ċ
ATOM	1424	OH	TYR	203	54. 259	74.586	22.970	1.00 7.05	A	0
ATOM	1425	C	TYR	203	55. 024	67. 234	24. 951	1.00 14.92	Ä	Č
ATOM	1425	0	TYR	203	53. 896	67. 124	25. 406	1.00 15.28	Ä	Ŏ
ATOM	1427	N	GLU	204	55. 744	66. 185	24. 570	1.00 16.35	Ä	Ň
ATOM	1421	CA	GLU	204	55. 222	64.826	24. 684	1.00 16.96	Ä	Ċ
			GLU	204	56. 238	63. 812	24. 130	1.00 14.28	Ä	Č
ATOM	1429	CB	GLU	204 204	55. 928	62.380	24. 130	1.00 14.20	A	č
ATOM	1430	CC	GLU	204 204	56. 872	61.345	23.947	1.00 14.51	A	č
ATOM	1431	CD	GLU	- 204 - 204	56. 697	60.144	24. 271	1.00 13.04	A	ŏ
ATOM	1432	0E1	GLU	204	50. 0 <i>51</i> 57. 778	61.714	23. 160	1.00 18.73	A	ő
ATOM	1433	OE2		204 204	54. 868	64. 431	26.128	1.00 18.02	A	Č
ATOM	1434	C	GLU	204 204	53.816	63.848	26. 388	1.00 18.02	A	Õ
ATOM	1435	0 M	GLU	204 205	55. 757	64. 761	27. 059	1.00 18.67	A	N
ATOM	1436	N	GLU		55. 589	64. 409	28. 459	1.00 10.01	A	C
ATOM	1437	CA	GLU	205	56. 970	64. 250	29. 096	1.00 20.30	A	Č
ATOM	1438	CB	GLU	205	56. 958	64. 035	30. 592	1.00 20.32	A	C
ATOM	1439	CC	GLU	205	56. 563	62.625	30. 974	1.00 24.02	A	Č
ATOM	1440	CD	GLU GLU	205	56. 398	62.355	32. 182	1.00 28.17	A	Ö
ATOM	1441	0E1		205		61.778	30.069	1.00 32.13	A	0
ATOM	1442		GLU	205	56.424		29. 319	1.00 31.11		C
ATOM	1443	C	GLU	205	54.760	65.362		1.00 22.23	A	
ATOM	1444	0	GLU	205	53.996	64.915	30. 164		A	0
ATOM	1445	N	GLU	206	54.902	66.666	29. 107	1.00 22.70	A	N
ATOM	1446	CA	GLU	206	54. 202	67.632	29. 939	1.00 23.19	A	C
ATOM	1447	CB	GLU	206	55. 203	68.667	30. 453	1.00 25.39	A	C
ATOM	1448	CG	GLU	206	56. 466	68.088	31.080	1.00 27.87	A	C
ATOM	1449	CD	GLU	206	56.188	67. 307	32. 345	1.00 29.45	A	C
ATOM	1450	0E1	GLU	206	57. 160	66.855	32.987	1.00 29.92	A	0
ATOM	1451	0E2	GLU	206	55.000	67.144	32.696	1.00 29.12	A	0
ATOM	1452	C	GLU	206	53. 024	68. 378	29. 324	1.00 24.91	A	C
ATOM	1453	0	GLU	206	52. 175	68. 885	30.051	1.00 24.03	A	0
ATOM	1454	И	VAL	207	52. 957	68.452	27. 999	1.00 25.41	A	N
ATOM	1455	CA	VAL	207	51.880	69. 199	27. 375	1.00 25.29	A	C
ATOM	1456	CB	VAL	207	52. 444	70. 235	26.398	1.00 25.95	A	C
ATOM	1457	CG1	VAL	207	51. 324	71.114	25.876	1.00 28.49	A	C
ATOM	1458		VAL	207	53. 496	71.080	27.092	1.00 26.77	A	C
ATOM	1459	C	VAL	207	50. 801	68. 409	26.653	1.00 26.09	A	C
ATOM	1460	0	VAL	207	49.617	68.703	26.813	1.00 27.62	A	0
ATOM	1461	N	PHE	208	51.194	67.412	25.865	1.00 26.41	A	N
ATOM	1462	CA	PHE	208	50. 228	66.620	25. 105	1.00 26.03	A	C
ATOM	1463	CB	PHE	208	50. 557	66.676	23.607	1.00 27.43	A	C
ATOM	1464	CG	PHE	208	50. 234	67.994	22.962	1.00 28.64	A	C
ATOM	1465		PHE	208	51.234	68.911	22. 679	1.00 29.07	A	C
ATOM	1466		PHE	208	48.918	68. 328	22.660	1.00 30.01	A	C
ATOM	1467		PHE	208	50.929	70. 142	22.104	1.00 30.28	A	C
ATOM	1468		PHE	208	48.604	69.556	22.086	1.00 30.23	A	C
ATOM	1469	CZ	PHE	208	49.612	70.464	21.809	1.00 30.40	A	C

					FΙ	G. 4	- 31			(Continued)
ATOM	1470	C	PHE	208	50. 082	65. 163	25. 506	1.00 26.13	A	C
ATOM	1471	0	PHE		49. 215	64. 471	24. 985	1.00 27.79	A	0
ATOM	1472	N	SER	209	50. 918	64. 687	26. 421	1.00 26.62	A	N
ATOM	1473	CA	SER	209	50. 852	63. 293	26. 848	1.00 25.74	A	C
ATOM	1474	CB	SER	209	49.645	63.059	27. 743	1.00 24.80	A	C
ATOM	1475	OG	SER	209	49.871	63. 629	29.014	1.00 29.47	A	0
ATOM ATOM	1476 1477	C	SER	209	50. 773	62.377	25.642	1.00 25.50	A	C
ATOM	1478	O N	SER ALA	209 210	50. 278 51. 272	61. 249 62. 875	25. 716 24. 524	1.00 25.72 1.00 23.72	A	0 N
ATOM	1479	CA	ALA	210	51. 263	62.112	23. 299	1.00 23.72	A	N C
ATOM	1480	CB	ALA	210	49. 977	62.364	23. 233	1.00 22.60	A A	Č
ATOM	1481	C	ALA	210	52. 455	62.560	22. 492	1.00 20.02	A	C
ATOM	1482	Õ	ALA	210	52. 986	63.644	22. 703	1.00 21.07	A	0
ATOM	1483	N	TYR	211	52. 863	61.719	21.558	1.00 21.57	A	N
ATOM	1484	CA	TYR	211	54. 000		20.718	1.00 21.42	Ä	Č
ATOM	1485	CB	TYR	211	54. 725	60.711	20. 405	1.00 19.58	A	Č ·
ATOM	1486	CG	TYR	211·	55. 921	60.870	19. 528	1.00 16.81	Ä	č
ATOM	1487		TYR	211	56. 853	61.870	19.770	1.00 16.07	Ä	č
ATOM	1488		TYR	211	58.002	61.971	19.001	1.00 18.18	Ä	Č
ATOM	1489	CD2	TYR	211	56. 160	59.976	18.489	1.00 17.91	Ā	Ċ
ATOM	1490	CE2	TYR	211	57. 306	60.065	17.716	1.00 18.80	Ā	Č
ATOM	1491	CZ	TYR	211	58. 221	61.063	17.979	1.00 18.36	Α	С
ATOM	1492	OH	TYR	211	59. 360	61.149	17.224	1.00 23.65	Α	0
ATOM	1493	C	TYR	211	53. 588	62.689	19.428	1.00 22.96	Α	C
ATOM	1494	0	TYR	211	54. 365	63.443	18.837	1.00 25.79	Α	0
ATOM	1495	N	SER	212	52. 365	62. 433	18.983	1.00 20.96	A	N
ATOM	1496	CA	SER	212	51.918	63.033	17.746	1.00 19.56	A	C
ATOM	1497	CB	SER	212	50. 835	62.175	17.090	1.00 20.97	A	C
ATOM	1498	OG	SER	212	49. 635	62. 208	17.829	1.00 21.79	A	0
MOTA	1499	C	SER	212	51.397	64. 439	17. 959	1.00 18.50	A	C
ATOM	1500	0 N	SER	212	50. 933	64. 789	19.040	1.00 16.31	A	0
ATOM ATOM	1501 1502	N	ALA	213	51. 493	65. 236	16.901	1.00 17.84	Ą	N .
ATOM	1502	CA CB	ALA ALA	$\begin{array}{c} 213 \\ 213 \end{array}$	51.036	66.610	16. 903	1.00 16.02	A	C
ATOM	1503	CP	ALA	213	52. 193 50. 429			1.00 14.16	A	C
ATOM	1505	Õ	ALA	213	50. 429	66. 935 67. 862	15. 526 14. 833	1.00 15.57 1.00 13.25	A	C
ATOM	1506	N	LEU	214	49. 448	66. 132	15. 129	1.00 13.25	A	0 N
ATOM	1507		LEU	214	48. 734	66. 339	13. 874	1.00 14.75	A A	N C
ATOM	1508		LEU	214	49. 353	65. 517	12. 735	1.00 16.03	A	Č
ATOM	1509		LEU	214	49. 482		12. 823	1.00 10.40	Ä	Č
ATOM	1510		LEU	214	48. 135	63.342	12.628	1.00 18.97	A	Č
ATOM	1511		LEU	214	50. 434		11.742	1.00 16.98	A	C C
ATOM	1512	C	LEU	214	47. 273	65.963	14. 124	1.00 16.65	A	Č
ATOM	1513		LEU	214	46. 966	64.933	14. 728	1.00 18.12	A	ŏ .
ATOM	1514	N	TRP	215	46. 366	66.811	13.666	1.00 16.16	Ä	N
ATOM	1515	CA	TRP	215	44.959	66. 590	13.907	1.00 14.69	Ä	Ċ
ATOM	1516	CB	TRP	215	44. 471	67.663	14.863	1.00 15.49	Ä	č
ATOM	1517	CG	TRP	215	45. 230	67.669	16.145	1.00 17.52	A	Č
ATOM	1518	CD2		215	46.482			1.00 17.74	Ä	č

					ास	G. 4	- 32			(Continued)
ATOM	1519	CRS	? TRP	215	46. 852	68.008	17. 729	1.00 17.50	A	C
ATOM	1520		TRP	215	47. 325	69. 149	15. 643	1.00 18.21	A	C
ATOM	1521		TRP	215	44. 904	67. 004	17. 289	1.00 15.79	A A	C C
ATOM	1522		TRP	215	45. 873	67. 202	18. 243	1.00 17.35	A	N
ATOM	1523		TRP	215	48. 033	68. 485	18. 318	1.00 18.06	A	Č
ATOM	1524		TRP	215	48. 505	69. 625	16. 228	1. 00 18. 96	A	Č .
ATOM	1525	CH2		215	48. 844	69. 289	17. 555	1. 00 18. 21	A	Č
ATOM	1526	C	TRP	215	44.110	66. 605	12.661	1.00 15.55	A	č
ATOM	1527	ŏ	TRP	215	43. 869	67.668	12. 090	1.00 16.18	A	ő
ATOM	1528	N	TRP	216	43.646	65. 430	12. 244	1.00 15.31	A	Ň
ATOM	1529	CA	TRP	216	42.793	65. 330		1.00 16.40	A	Ċ
ATOM	1530	CB	TRP	216	42.494	63.873	10.739	1.00 16.43	Ä	Č
ATOM	1531	CG	TRP	216	43. 549	63. 114	10.002	1.00 17.38	Ā	Č
ATOM	1532	CD2	TRP	216	43. 823	63.169	8. 599	1.00 17.01	A	Ċ
ATOM	1533		TRP	216	44. 794	62.176	8.320	1.00 17.25	Α	Ċ
ATOM	1534		TRP	216	43.340	63.954	7. 549	1.00 17.09	Α	C
ATOM	1535		TRP	216	44.352	62. 125	10. 508	1.00 18.55	Α	C
ATOM	1536		TRP	216	45.098	61.553	9. 501	1.00 18.07	Α	N
ATOM	1537		TRP	216	45. 286	61.951	7.036	1.00 15.24	Α	C
ATOM	1538		TRP	216	43.829	63. 729	6. 270	1.00 17.06	A	C
ATOM	1539	CH2		216	44. 794	62. 734	6.027	1.00 17.07	A	Ċ
ATOM	1540	C	TRP	216	41.461	66.016	11. 355	1.00 17.17	A	C
ATOM	1541	0	TRP	216	40.990	66.005	12. 487	1.00 18.00	A	0
ATOM ATOM	1542 1543	N Ca	SER SER	$\begin{array}{c} 217 \\ 217 \end{array}$	40.847	66.605	10. 334	1.00 18.39	A	N
ATOM	1544	CB	SER	217	39. 552 39. 257	67. 240 68. 225	10. 523	1.00 19.62	A	C C
ATOM	1545	OG	SER	217	39. 234	67. 589	9. 392 8. 133	1.00 20.31 1.00 24.00	A	
ATOM	1546	C	SER	217	38. 528	66. 108	10. 550	1.00 24.00	A A	0 C
ATOM	1547	Õ	SER	217	38. 814	64. 994	10. 110	1.00 20.47	A	0
ATOM	1548	Ň	PRO	218	37. 326	66.369	11.074	1.00 20.32	A	N N
ATOM	1549	CD	PRO	218	36. 827	67.650	11.598	1.00 20.28	A	Č
ATOM	1550	CA	PRO	218	36. 285	65. 339	11. 154	1.00 22.67	A	Č.
ATOM	1551	CB	PRO	218	35. 033	66. 148	11.462	1.00 21.68	Ä	č
ATOM	1552	CG	PRO	218	35.587	67. 223	12.353	1.00 21.12	Ä	č
ATOM	1553	C	PRO	218	36.123	64.404	9.950	1.00 23.46	A	Č
ATOM	1554	0	PRO	218	36.190	63.183	10.107	1.00 25.13	A	0
ATOM	1555	N	ASN	219	35.909	64. 948	8.756	1.00 22.93	A	N
ATOM	1556	CA	ASN	219	35.756	64.071	7.600	1.00 22.31	A	C
ATOM	1557	CB	ASN	219	34. 704	64.622	6.631	1.00 22.48	Α	C
ATOM	1558	CG	ASN	219	35. 172	65.849	5. 903	1.00 24.12	Α	C
ATOM	1559		ASN	219	36. 373	66.076	5.760	1.00 26.01	Α	0
ATOM	1560		ASN	219	34. 230	66.640	5. 411	1.00 26.27	A	N
ATOM	1561	C	ASN	219	37. 090	63. 841	6.871	1.00 21.20	A	C
ATOM	1562	0 M	ASN	219	37. 115	63. 307	5. 760	1.00 20.94	A	0
ATOM	1563	N	GLY	220	38. 184	64. 267		1.00 18.33	A	N
ATOM	1564	CA	GLY	220	39. 512	64.068	6. 941	1.00 17.97	A	C
ATOM	1565 1566	C 0	GLY GLY	220 220	40.035	64. 993	5.853	1.00 18.92	A	C
ATOM	1567	N	THR	220 221	41. 157	64. 801	5.375	1.00 20.28	· A	0
ATOM	1001	17	TIM	441	39. 242	65.980	5.447	1.00 17.57	A	N

										(Continued)
				-	F	I G. 4	- 34			(00221223334)
ATOM	1617	СВ	GLN	227 .	56.09	6 73.423	3 13. 331	1.00 15.62	A	С
ATOM	1618	CG	GLN	227	57. 51	4 73.24	6 12.799	1.00 16.35	A	C
ATOM	1619	CD	GLN	227	57.84	7 74.19	11.666	1.00 14.31	A	C
ATOM	1620	0E1	GLN	227	57.87			1.00 18.11		0
ATOM	1621	NE2		227	58. 10			1.00 12.45		N
ATOM	1622	C	GLN	227	56.61			1.00 16.27		C
ATOM	1623	0	GLN	227	56.34			1.00 16.33		0
ATOM	1624	N	PHE	228	57.60			1.00 17.36		N
ATOM	1625	CA	PHE	228	58. 41			1.00 16.81		C
ATOM	1626	CB	PHE	228	58. 32			1.00 14.62		C
ATOM	1627	CG	PHE	228	56.91			1.00 14.48		C
ATOM	1628	CD1		228	56. 31			1.00 14.37		C
ATOM	1629	CD2		228	56.18			1.00 12.73		C
ATOM	1630	CE1		228	55. 00			1.00 13.56		C
ATOM	1631		PHE	228	54. 87 54. 27			1.00 14.73		C
ATOM ATOM	1632 1633	CZ C	PHE PHE	228 228	59. 84			1.00 15.31 1.00 18.12		C
ATOM	1634	0	PHE	228	60.41			1.00 17.47		ŏ
ATOM	1635	N	ASN	229	60.41			1.00 20.00		N
ATOM	1636	CA	ASN	229	61.77			1.00 20.87		Č
ATOM	1637	CB	ASN	229	61.76			1.00 21.57		č
ATOM	1638		ASN	229	63.08			1.00 24.35		Č
ATOM	1639		ASN	229	64.14			1.00 26.00		0
ATOM	1640		ASN	229	63.02			1.00 25.62		N
ATOM	1641	C	ASN	229	62.54	0 74.42	18.362	1.00 21.39	A	C
ATOM	1642	0	ASN	229	62. 23			1.00 21.52		0
ATOM	1643	N	ASP	230	63.51			1.00 20.96		N
ATOM	1644	CA	ASP	230	64. 30			1.00 22.78		C
ATOM	1645	CB	ASP	230	64. 27			1.00 22.69		C
ATOM	1646	CG	ASP	230	62.88			1.00 22.37		C
ATOM	1647		ASP	230	62. 68			1.00 21.57		0
ATOM	1648		ASP	230	61.99			1.00 21.82		0
ATOM ATOM	1649 1650	C 0	ASP ASP	230 230	65. 73 66. 66			1.00 24.50 1.00 24.72		C 0
ATOM	1651	N	THR	231	65. 90			1.00 24.72		N N
ATOM	1652	CA	THR	231	67. 22			1.00 26.22		Č
ATOM	1653	CB	THR	231	67. 14			1.00 27.87		Č
ATOM	1654	OG1		231	66. 54			1.00 28.62		ŏ
ATOM	1655		THR	231	68. 54			1.00 26.63		č
ATOM	1656	C	THR	231	68. 09			1.00 26.77		Č
ATOM	1657	Ŏ	THR	231	69. 25			1.00 27.34		Ö
ATOM	1658	N	GLU	232	67.55			1.00 25.50		N
ATOM	1659	CA	GLU	232	68. 32	9 77.02		1.00 24.52		C
ATOM	1660	CB	GLU	232	68.15			1.00 28.36		C
ATOM	1661	CG	GLU	232	68. 61			1.00 34.72		C
ATOM	1662	CD	GLU	232	68. 48			1.00 40.02		C
ATOM	1663	OE1		232	68. 76			1.00 44.21		0
ATOM	1664		GLU	232	68.10			1.00 42.26		0
ATOM	1665	С	GLU	232	68.02	0 76.37	7 22.627	1.00 22.97	' A	С

(Continued)

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FIG. 4-35 **ATOM GLU** 232 76.942 23.679 1666 0 68.331 1.00 20.81 0 ATOM 1667 N VAL 233 67.416 75.194 22.596 1.00 20.32 N Α **ATOM** 1668 CA VAL 233 67.091 74.499 23.832 1.00 17.88 C Α 233 73.618 23.648 C 1669 CB VAL 65.853 1.00 17.88 **ATOM** Α C **ATOM** 1670 CG1 VAL 233 65.522 72.925 24.957 1.00 14.00 A 233 Ċ 74.478 **ATOM** 1671 CG2 VAL 64.678 23.160 1.00 16.73 A 233 C **ATOM** 1672 VAL 68. 261 73.642 24.304 C 1.00 16.00 Α ATOM 1673 0 VAL 233 68.694 72.728 23.606 1.00 15.94 0 A 1674 N 234 73.927 25.504 N ATOM PR₀ 68.788 1.00 14.51 Α 234 74.907 C **ATOM** 1675 CD PR₀ 68.313 26.494 1.00 13.03 A C 1676 CA 234 69.914 73.162 26.040 1.00 13.93 ATOM PR₀ A C CB 234 70.031 73.677 27.473 ATOM 1677 **PRO** 1.00 12.63 A 1678 **PRO** 234 69.517 75.059 27.377 C ATOM CG 1.00 11.32 Α ATOM 1679 C PR₀ 234 69.643 71.663 25.987 1.00 16.20 A **ATOM** 1680 PR₀ 234 68.487 71.220 26.041 0 0 1.00 15.73 Α 1681 70.716 N 235 70.887 25.900 N ATOM LEU 1.00 16.28 A 1682 CA LEU 235 70.602 69.443 25.825 ATOM 1.00 16.91 A ${\rm C}$ CB ATOM 1683 LEU 235 71.505 68.912 24.718 1.00 18.54 A 235 **ATOM** 1684 CG LEU 71.267 69.349 23.273 1.00 21.93 $\begin{array}{c} C \\ C \\ C \\ C \end{array}$ A 235 **ATOM** 1685 CD1 LEU 72.434 68.856 22.412 1.00 21.90 A **ATOM** 1686 CD2 LEU 235 69.946 68.790 22.768 1.00 19.17 A **ATOM** 1687 LEU 235 70.990 68.743 C 27.118 1.00 17.26 A **ATOM** 1688 235 71.939 LEU 69.157 27.793 0 0 1.00 18.36 A 1689 ILE 70. 244 **ATOM** 67.696 27.472 N 236 1.00 14.95 N A ATOM 1690 CA ILE 236 70.586 66.899 28.644 1.00 12.68 A C CB **ATOM** 1691 ILE 236 69.345 66.245 29.335 1.00 10.50 A **ATOM** 1692 CG2 ILE 236 68.538 65.433 28.329 1.00 9.32 C A CG1 ILE 236 **ATOM** 1693 69.806 65.298 30.448 1.00 8.74 A **ATOM** 1694 CD1 ILE 236 70.789 65.919 31.427 C 1.00 7.11 A ATOM 1695 C ILE 236 71.444 65.802 28.010 1.00 12.84 A **ATOM** 1696 0 ILE 236 71.105 65.276 26.942 0 1.00 10.11 A 1697 GLU **ATOM** N 237 72.558 65.480 28.650 1.00 12.44 N C C A ATOM 1698 CA **GLU** 237 64.470 73.463 28.128 1.00 14.46 A CB 237 **ATOM** 1699 **GLU** 74.767 65.128 27.655 1.00 13.45 A CG 237 **ATOM** 1700 **GLU** 74.554 66.079 26.500 1.00 18.02 C A 237 CD **ATOM** 1701 **GLU** 75.845 66.500 25.819 1.00 23.46 A ATOM 1702 OE1 GLU 237 0 75.779 67.016 24.683 1.00 25.80 A **ATOM** 1703 OE2 GLU 237 76.928 66.324 26.408 1.00 26.23 0 A 237 **ATOM** 1704 GLU 73.744 C C 63.427 29.191 1.00 13.41 A 1705 237 **ATOM** 0 **GLU** 73.895 63.752 30.363 1.00 14.43 A 0 238 1706 N 62.169 ATOM TYR 73.801 28.781 1.00 12.83 A N **ATOM** 1707 CA TYR 238 74.052 61.093 29.721 1.00 14.06 A C 238 Č **ATOM** 1708 CB TYR 72.810 60.840 30.595 1.00 12.42 A 1709 CG 238 **ATOM** TYR 71.566 60.419 29.856 1.00 11.79 $_{\rm C}^{\rm C}$ A 238 **ATOM** 1710 CD1 TYR 71.451 59.139 29.317 1.00 16.12 A CE1 TYR 238 70. 292 ATOM 1711 58.739 28.635 1.00 17.09 ${\bf C}$ A TYR ATOM 1712 CD2 238 70.496 61.295 29.701 1.00 12.13 A **ATOM** 1713 CE2 TYR 238 69.336 60.913 29.020 1.00 12.94 ${\bf C}$ A CZ238 **ATOM** 1714 TYR 69.243 59.634 28.487 1.00 15.48

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	• •	F	IG. 4-36		(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1715 OH TYR 1716 C TYR 1717 O TYR 1718 N SER 1719 CA SER 1720 CB SER 1721 OG SER 1722 C SER 1723 O SER 1724 N PHE 1725 CA PHE 1726 CB PHE 1727 CG PHE 1728 CD1 PHE 1729 CD2 PHE 1730 CE1 PHE 1731 CE2 PHE 1731 CE2 PHE	238 68. 12 238 74. 44 238 74. 05 239 75. 22 239 75. 68 239 76. 92 239 77. 90 239 74. 66 239 73. 75 240 74. 80 240 73. 97 240 71. 89 240 70. 82 240 71. 98 240 71. 98 240 71. 98 240 71. 98 240 71. 98 240 71. 91 240 69. 85 240 71. 01 240 69. 95	27 59. 257 27. 775 15 59. 847 28. 954 19 59. 667 27. 798 20 58. 986 29. 596 39 57. 779 28. 943 26 57. 251 29. 656 20 58. 265 29. 766 31 56. 668 28. 879 35 56. 587 29. 700 39 55. 834 27. 862 22 54. 678 27. 679 33 54. 833 26. 523 36 53. 843 26. 574 24 54. 037 27. 436 30 52. 655 25. 858 39 53. 064 27. 597 8 51. 675 26. 012 54 51. 878 26. 888	1. 00 15. 25 1. 00 17. 74 1. 00 14. 10 1. 00 13. 87 1. 00 11. 90 1. 00 18. 76 1. 00 13. 45 1. 00 14. 39 1. 00 12. 12 1. 00 12. 12 1. 00 12. 48 1. 00 11. 50 1. 00 10. 15 1. 00 10. 78 1. 00 11. 03 1. 00 10. 46	A
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1733 C PHE 1734 O PHE 1735 N TYR 1736 CA TYR 1737 CB TYR 1738 CG TYR 1739 CD1 TYR 1740 CE1 TYR 1741 CD2 TYR 1742 CE2 TYR 1742 CE2 TYR 1743 CZ TYR 1744 OH TYR 1745 C TYR 1746 O TYR 1747 N SER 1748 CA SER 1749 CB SER	240 75. 01 240 75. 72 241 75. 12 241 76. 14 241 76. 52 241 76. 83 241 78. 32 241 75. 86 241 76. 10 241 77. 33 241 75. 79 241 76. 68 242 74. 50 242 74. 46 242 74. 46	8 53. 652 27. 330 22 53. 805 26. 335 29 52. 617 28. 153 47 51. 612 27. 958 26 51. 057 29. 329 23 52. 167 30. 317 25 52. 821 30. 308 26 53. 894 31. 168 26 53. 678 32. 080 38 54. 319 32. 046 36 55. 408 32. 859 33 50. 510 26. 967 36 49. 948 26. 322 31 50. 204 26. 837 33 49. 180 25. 888 34 49. 590 24. 469	1. 00 14. 83 1. 00 18. 18 1. 00 13. 74 1. 00 13. 29 1. 00 13. 69 1. 00 10. 88 1. 00 11. 93 1. 00 9. 47 1. 00 12. 15 1. 00 11. 02 1. 00 12. 15 1. 00 10. 38 1. 00 14. 62 1. 00 12. 20 1. 00 16. 13 1. 00 16. 13 1. 00 16. 30	A C A C A C A C A C A C A C A C A C A C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1750 OG SER 1751 C SER 1752 O SER 1753 N ASP 1754 CA ASP 1755 CB ASP 1756 CG ASP 1757 OD1 ASP 1758 OD2 ASP 1759 C ASP 1760 O ASP 1761 N GLU 1762 CA GLU 1763 CB GLU	242 74.64 242 75.21 243 74.51 243 75.06 243 74.77 243 73.29 243 72.86 243 76.57 243 77.29 244 77.01 244 78.41	47 47. 816 26. 226 19 47. 625 27. 303 16 46. 865 25. 312 36 45. 535 25. 548 74 44. 605 24. 369 30 44. 419 24. 132 49 44. 246 25. 126 32 44. 438 22. 955 72 45. 554 25. 805 38 46. 432 25. 330 46 43. 559 26. 567 42 44. 363 26. 944	1. 00 17. 46 1. 00 19. 13 1. 00 19. 34 1. 00 23. 36 1. 00 27. 30 1. 00 33. 83 1. 00 36. 97 1. 00 37. 15 1. 00 23. 56 1. 00 22. 48 1. 00 24. 45 1. 00 22. 80	A 0 A C A 0 A N A C A C A 0 A 0 A 0 A 0 A 0

					FΙ	G. 4	- 37			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1764 1765 1766 1767 1768 1769 1770 1771 1772 1773 1774 1775 1776 1777 1778	OE2 C O N CA CB OG C O N CA CB CC CD1	GLU GLU GLU GLU GLU SER SER SER SER LEU LEU LEU	244 244 244 244 245 245 245 245 246 246 246 246	79. 940 79. 967 81. 079 78. 877 79. 374 80. 533 78. 888 79. 724 79. 080 77. 949 80. 044 80. 874 79. 392 79. 694 78. 522 78. 659 78. 736	42. 547 41. 177 40. 680 40. 601 44. 476 44. 854 44. 159 44. 205 43. 402 44. 068 45. 605 45. 762 46. 628 47. 983 48. 926 50. 368 50. 388	27. 995 28. 667 28. 958 28. 903 25. 754 25. 913 24. 561 23. 370 22. 244 21. 723 22. 861 21. 971 23. 397 22. 943 23. 229 22. 728 21. 214	1. 00 29. 35 1. 00 29. 80 1. 00 29. 53 1. 00 29. 32 1. 00 22. 28 1. 00 21. 94 1. 00 19. 92 1. 00 19. 31 1. 00 17. 93 1. 00 18. 69 1. 00 18. 41 1. 00 18. 20 1. 00 17. 99 1. 00 16. 83	A A A A A A A A A A A A A A A A A A A	C C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1781 1782 1783 1784 1785 1786 1787 1788 1789 1790 1791 1792 1793 1794	CD2 C O N CA CB CG CD OE1	LEU LEU LEU GLN GLN GLN GLN GLN GLN TYR	246 246 246 247 247 247 247 247 247 247 247 247 248 248	77. 458 80. 943 80. 921 82. 034 83. 295 84. 400 85. 791 86. 875 86. 829 87. 862 83. 224 83. 640 82. 710 82. 592	51. 181 48. 463 48. 662 48. 635 49. 073 49. 038 49. 234 48. 770 49. 065 48. 049 50. 461 50. 648 51. 436 52. 794	23. 192 23. 679 24. 895 22. 940 23. 532 22. 480 23. 045 22. 090 20. 899 22. 611 24. 170 25. 313 23. 430 23. 954	1.00 10.83 1.00 19.98 1.00 18.12 1.00 16.81 1.00 17.30 1.00 15.11 1.00 17.62 1.00 18.47 1.00 20.53 1.00 17.76 1.00 17.66 1.00 17.56 1.00 18.50 1.00 19.00	A A A A A A A A A A A A	C C O N C C C C O .N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1795 1796 1797 1798 1799 1800 1801 1802 1803 1804 1805 1806 1807 1808 1809 1810 1811 1812	CE1 CD2	TYR	248 248 248 248 248 248 248 248 248 249 249 249 249 249 249 250	83. 177 84. 684 85. 353 86. 742 85. 444 86. 839 87. 479 88. 854 81. 130 80. 288 80. 804 81. 610 79. 411 79. 424 80. 857 78. 937 79. 734 77. 638	53. 822 53. 820 52. 812 52. 814 54. 838 54. 851 53. 836 53. 809 53. 134 53. 549 53. 595 53. 886 54. 222 54. 582 55. 042 55. 042 55. 864 55. 096	22. 972 22. 860 22. 172 22. 058 23. 437 23. 333 22. 647 22. 595 24. 212 23. 323 25. 440 26. 668 27. 206 27. 206 27. 481 24. 852 24. 413 24. 599	1. 00 17. 39 1. 00 16. 80 1. 00 17. 20 1. 00 17. 58 1. 00 17. 77 1. 00 17. 22 1. 00 18. 42 1. 00 19. 27 1. 00 18. 87 1. 00 19. 15 1. 00 18. 20 1. 00 18. 21 1. 00 18. 83 1. 00 19. 46 1. 00 17. 63 1. 00 19. 66 1. 00 20. 92 1. 00 19. 01	A A A A A A A A A A A A A A A A A A A	C C C C C C C C C C C C C C C C C C C

(Continued) FIG. 4-38 C 56.158 23. 785 1,00 19,61 77.083 1813 CA LYS 250 ATOM C 55.618 22.936 1.00 23.51 75.933 1814 CB LYS 250 Α **ATOM** C 1.00 28.40 CG 76.320 54. 428 22.089 Α LYS 250 **ATOM** 1815 C 75.197 54.010 21.152 1.00 30.62 A 250 **ATOM** 1816 CD LYS 52.938 20. 203 1.00 32.02 C 75.698 A 250 CE LYS **ATOM** 1817 53. 385 19.546 1.00 32.62 N 76.966 A 1818 NZ LYS 250 ATOM C 24.628 1.00 17.92 76.580 57.320 Α 250 ATOM 1819 C LYS 25.758 0 1.00 17.90 Α 76.130 57.130 **ATOM** 1820 0 LYS 250 N 76.663 58.524 24.077 1.00 14.61 A 1821 N THR 251 ATOM 59.689 24.786 1.00 15.48 C 76.171 Α 1822 CA THR 251 **ATOM** C 60.887 24.666 1.00 13.61 A 77.104 CB 251 **ATOM** 1823 THR 0 25.441 1.00 15.96 78.280 60.654 Α ATOM 1824 OG1 THR 251 25.181 C 76.414 62.137 1.00 13.93 CG2 THR 251 Α **ATOM** 1825 24. 205 C 74.832 60.086 1.00 16.04 Α 1826 C THR 251 **ATOM** 74.755 23.083 1.00 17.34 Α 0 60.572 **ATOM** 1827 0 THR 251 73.779 24.977 N 1828 59.860 1.00 15.27 Α VAL 252 ATOM N 24.559 C 72.439 60.205 1.00 16.08 VAL Α 1829 CA 252 **ATOM** C 59.381 25.355 1.00 16.76 71.405 Α 1830 CB VAL 252 **ATOM** 69.987 59.832 25.014 1.00 16.29 Α C 1831 CG1 VAL 252 **ATOM** 25.050 1832 71.595 57.895 1.00 13.65 Α ATOM CG2 VAL 252 1833 72.223 61.699 24.799 1.00 18.46 C 252 Α **ATOM** VAL 72.443 62.212 25.905 1.00 19.01 A 0 1834 0 VAL 252 ATOM 62.398 23.754 1.00 19.18 A 71.799 1835 N 253 **ATOM** ARG 23.842 1836 253 71.568 63.831 1.00 18.54 Α **ATOM** CA ARG 22.949 72.574 1.00 19.46 C 1837 CB ARG 253 64.567 Α **ATOM** 1.00 24.49 C ATOM 74.014 64.439 23.457 A 1838 CG ARG 253 65.066 22.519 1.00 29.04 1839 CD ARG 253 75.021 A ATOM 1840 253 75.797 64.044 21.822 1.00 35.89 N **ATOM** NE ARG A 77.013 63.647 22.185 1.00 38.08 1841 CZC 253 ATOM ARG 1.00 39.69 23.241 77.606 64.191 A N ATOM 1842 NH1 ARG 253 21.497 1843 NH2 ARG 77.633 62.699 1.00 40.12 N **ATOM** 253 70.140 64.156 23.449 1.00 17.33 C **ATOM** 1844 C **ARG** 253 A 22.362 0 69.690 63.802 1.00 18.44 0 **ATOM** 1845 ARG 253 Α 69.432 64.836 24.344 1.00 16.85 N 1846 N VAL 254 A **ATOM** 24.125 65.196 1.00 15.67 CA 68.033 A **ATOM** 1847 VAL 254 67.079 64.405 25.070 1.00 16.67 **ATOM** 1848 CB VAL 254 A 24.766 C 1849 254 65.640 64.775 1.00 16.79 Α **ATOM** CG1 VAL CG2 VAL 67.308 24.951 C 254 62.899 1.00 17.24 Α **ATOM** 1850 C 1851 C 254 67.737 66.660 24.405 1.00 14.62 Α **ATOM** VAL 1852 0 254 68.122 67.186 25.450 1.00 15.12 0 **ATOM** VAL Α 67.048 23.475 1.00 13.71 N **ATOM** 1853 N PR₀ 255 67.340 Α **ATOM** CD 66.677 66.945 22.105 1.00 10.62 Α C 1854 PR₀ 255 C 66.725 68.749 23.730 1.00 13.00 1855 CA PR₀ 255 A ATOM 66.064 69.193 22.431 1.00 13.28 C 1856 CB PR₀ 255 A **ATOM** C 21.397 PR₀ 255 66.674 68.265 1.00 13.45 Α 1857 CG **ATOM** 24.899 1.00 13.86 C 255 68.674 **ATOM** 1858 C PR₀ 65.735 0 ATOM 1859 0 PR₀ 255 64.663 68.086 24.772 1.00 13.58 1860 256 66.108 69.255 26.032 1.00 13.63 N N TYR ATOM

SUBSTITUTE SHEET (RULE 26)

69.194

27. 242

65.304

256

CA

1861

ATOM

TYR

1.00 11.65

						_				(Continued)
					FΙ	G. 4	- 39			
ATOM ATOM ATOM	1862 1863 1864	CB CG	TYR TYR TYR	256 256 256	65. 801 65. 044 64. 949	68. 006 67. 706 68. 646	28. 077 29. 351 30. 378	1.00 10.57 1.00 10.49 1.00 9.61	A A A	C C C
ATOM ATOM	1865 1866	CE1	TYR TYR	256 256	64. 296 64. 460	68. 351 66. 460	31. 571 29. 549	1.00 7.54 1.00 9.65	A A	C C
ATOM ATOM	1867 1868	CZ	TYR TYR	256 256	63. 799 63. 722	66. 156 67. 105	30. 735 31. 742	1.00 11.05 1.00 10.10	A A	C C
ATOM ATOM ATOM	1869 1870 1871	OH OH	TYR TYR TYR	256 · 256 256	63. 060 65. 488 66. 559	66. 801 70. 492 70. 750	32. 909 28. 012 28. 553	1.00 10.49 1.00 12.70 1.00 15.49	. A . A . A	0 C 0
ATOM ATOM	1872 1873	N CD	PRO PRO	257 257	64. 444 63. 174	71. 325 71. 254	28. 080 27. 334	1.00 13.45 1.00 12.39 1.00 13.82	A A A	N C
ATOM ATOM	1874 1875	CA CB	PRO PRO	257 257	64. 548 63. 501	72. 593 73. 450	28. 800 28. 106	1.00 11.47 1.00 12.01	A A	C C
ATOM ATOM ATOM	1876 1877 1878	CG C O	PRO PRO PRO	257 257 257	62. 405 64. 296 63. 174	72. 464 72. 489 72. 210	27. 866 30. 298 30. 723	1.00 12.87 1.00 12.85 1.00 15.59	· A A A	C C O
ATOM ATOM	1879 1880	N CA	LYS LYS	258 258	65. 327 65. 155	72. 718 72. 671	31.105 32.546	1.00 11.64 1.00 11.10	A A	N C
ATOM ATOM ATOM	1881 1882 1883	CB CG CD	LYS LYS LYS	258 258 258	66. 501 67. 034 68. 519	72. 439 71. 012 70. 906	33. 227 33. 031 33. 331	1. 00 12. 96 1. 00 14. 20 1. 00 13. 34	A A	C C C
ATOM ATOM	1884 1885	CE NZ	LYS LYS	258 258	69. 042 68. 671	69. 480 68. 536	33. 136 34. 223	1.00 13.95 1.00 10.80	A A A	C N
ATOM ATOM ATOM	1886 1887 1888	C O N	LYS LYS ALA	258 258 259	64. 517 64. 368	73. 984 74. 921	33. 011 32. 224	1.00 12.44	A A	C 0
ATOM ATOM	1889 1890	CA CB	ALA ALA	259 259 259	64. 124 63. 484 63. 368	74. 043 75. 236 75. 097	34. 280 34. 844 36. 355	1. 00 13. 33 1. 00 14. 81 1. 00 16. 40	A A · A	N C C
ATOM ATOM	1891 1892	C 0	ALA ALA	259 259	64. 167 65. 317	76. 555 76. 787	34. 508 34. 881	1.00 15.14 1.00 17.32	A A	C 0
ATOM ATOM ATOM	1893 1894 1895	N CA C	GLY GLY GLY	260 260 260	63. 448 63. 984 64. 870	77. 419 78. 720 78. 749	33. 802 33. 444 32. 217	1.00 16.82 1.00 15.59 1.00 15.78	A A A	N C C
ATOM ATOM	1896 1897	0 N	GLY ALA	260 261	65. 379 65. 072	79. 812 77. 600	31.852 31.577	1.00 17.65 1.00 13.77	A A	O N
ATOM ATOM ATOM	1898 1899 1900	CA CB C	ALA ALA ALA	261 261 261	65. 906 66. 524 65. 093	77. 554 76. 182 77. 911	30. 379 30. 224 29. 137	1.00 11.19 1.00 10.21 1.00 10.04	A A A	C C C
ATOM ATOM	1901 1902	0 N	ALA VAL	261 262	63. 896 65. 747	78. 160 77. 947	29. 212 27. 987	1.00 10.04 1.00 8.71 1.00 11.73	A A	O N
ATOM ATOM ATOM	1903 1904 1905	CA CB CG1	VAL VAL	262 262 262	65.050 66.035 65.257	78. 284 78. 529 78. 796	26. 761 25. 594 24. 299	1.00 12.13 1.00 11.50	A A	C ·
ATOM ATOM ATOM	1906 1907	CG2 C		262 262 262	66. 939 64. 092	79. 732 77. 167	24. 299 25. 920 26. 389	1.00 8.31 1.00 5.79 1.00 13.92	A A A	C C C
ATOM ATOM	1908 1909	O N	VAL ASN	262 263	64. 471 62. 844	76. 001 77. 536	26. 341 26. 139	1.00 16.73 1.00 13.49	A A	O N
ATOM	1910	CA	ASN	263	61.816	76. 585	25. 773	1.00 13.67	A	С

										(Cont	inued)
					FΙ	G. 4	- 40			,	
ATOM	1911	СВ	ASN	263	60. 470	77. 038	26. 336	1.00 14.53	Α	С	
ATOM	1912	CG	ASN	263	60. 222		27.746	1.00 17.27	Ā	Č	
ATOM	1913		ASN	263	59. 342		28. 444	1.00 18.62	Ā	Õ	
ATOM	1914		ASN	263	60.977		28.169	1.00 16.78	Ā	N	
ATOM	1915	C	ASN	263	61.715		24.265	1.00 14.45	A	C	
ATOM	1916	Ŏ	ASN	263	62. 170		23.561	1.00 16.33	Ā	0	
ATOM	1917	Ň	PRO	264	61.119		23.743	1.00 14.86	Α	N	
ATOM	1918	CD	PRO	264	60.513		24.412	1.00 15.86	Α	C	
ATOM	1919	CA	PRO	264	60. 986		22. 294	1.00 15.41	Α	С	
ATOM	1920	CB	PR0	264	60. 591	73.844	22.106	1.00 14.97	Α	C	
ATOM	1921	CG	PRO	264	59. 721	73.607	23. 287	1.00 14.81	Α	C	
ATOM	1922	C	PR0	264	59.867		21.882	1.00 15.66	Α	C	
ATOM	1923	0	PR0	264	58.954		22.663	1.00 17.42	Α	0	
ATOM	1924	N	THR	265	59. 942		20.673	1.00 15.76	Α	N	
ATOM	1925	CA	THR	265	58. 895		20. 199	1.00 14.67	Å	C	
ATOM	1926	CB	THR	265	59. 458		19.341	1.00 15.37	Α	C	
ATOM	1927		THR	265	60. 162		18. 223	1.00 15.98	A	0	
ATOM	1928		THR	265	60. 402		20. 159	1.00 12.01	A	C	
ATOM	1929	C	THR	265	58. 024		19.360		. A	C	
ATOM	1930	0	THR	265	58. 465		18. 932	1.00 18.75	A	0	
ATOM	1931	N	VAL	266	56. 794		19. 113	1.00 15.56	A	N	
ATOM	1932	CA	VAL	266	55. 872		18. 347	1.00 12.79	A	C	
ATOM	1933	CB	VAL	266	54. 856		19. 274	1.00 12.90	A	C	
ATOM	1934		VAL	266	54. 193		20.130	1.00 12.06	A	C	
ATOM	1935		VAL	266	53. 821		18.466	1.00 10.69	A	C	
ATOM	1936	C	VAL	266	55. 115		17.350	1.00 12.88	A	C	
ATOM	1937	0 N	VAL LYS	$\begin{array}{c} 266 \\ 267 \end{array}$	54. 995		17. 511 16. 327	1.00 12.12	A	0	
ATOM ATOM	1938 1939	N CA	LYS	267	54. 601 53. 817		15. 262	1.00 13.52 1.00 13.08	A	N C	
ATOM	1939	CB	LYS	267	54. 692		14. 050	1.00 13.06	A A	C	
ATOM	1941	CG	LYS	267	55. 642		14. 165	1.00 13.04	A	C	
ATOM	1942	CD	LYS	267	56. 348		12. 833	1.00 11.33	A	Č	
ATOM	1943	CE	LYS	267	57. 313	79.864	12. 788	1.00 11.66	A	C	
ATOM	1944	NZ	LYS	267	58. 007	79.844	11. 459	1.00 12.98	Ä	N	
ATOM	1945			267	52. 713			1.00 14.81	Ä		
ATOM	1946	ŏ	LYS	267	52. 885		14. 930	1.00 14.91	· A	ő	
ATOM	1947	Ň	PHE	268	51.588		14. 389	1.00 15.02	Ä	Ň	
ATOM	1948	ĊA	PHE	268	50. 471		13. 975	1.00 14.84	Ä	Ĉ	
ATOM	1949	CB	PHE	268	49. 249		14.842	1.00 13.98	Ä	Č	
ATOM	1950	CG	PHE	268	48. 237		14.846	1.00 15.65	Ä	Č	
ATOM	1951		PHE	268	48. 467		15.562	1.00 15.51	Ä	Č	
ATOM	1952		PHE	268	47.056		14. 115	1.00 18.05	Ä	Č	
ATOM	1953		PHE	268	47. 537		15.551	1.00 15.17	A	Č	
ATOM	1954		PHE	268	46.120		14.101	1.00 17.28	A	C	
ATOM	1955	CZ	PHE	268	46. 366		14.821	1.00 14.54	Ā	Ċ	
ATOM	1956	С	PHE	268	50.117		12.497	1.00 14.63	Α	Ç.	
ATOM	1957	0	PHE	268	50.143		11.981	1.00 16.53	Α	0	
ATOM	1958	N	PHE	269	49.767	74.938	11.829	1.00 13.37	A	N	
ATOM	1959	CA	PHE	269	49.417	74.976	10.413	1.00 12.73	Α	С	

					FIG. 4-41	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1976	CD2 CE2 CZ C C O N CA CB CG1	PHE PHE PHE PHE PHE PHE VAL VAL VAL VAL VAL VAL VAL	269 269 269 269 269 269 269 270 270 270 270 270 270 270 271 271	50. 597 74. 510 9. 547 1. 00 12. 68 A 51. 875 75. 229 9. 809 1. 00 10. 71 A 52. 190 76. 387 9. 112 1. 00 11. 11 A 52. 758 74. 759 10. 770 1. 00 11. 04 A 53. 374 77. 070 9. 371 1. 00 12. 54 A 53. 940 75. 430 11. 039 1. 00 13. 96 A 54. 252 76. 591 10. 339 1. 00 13. 89 A 48. 270 74. 032 10. 117 1. 00 12. 37 A 47. 937 73. 157 10. 910 1. 00 14. 50 A 47. 699 74. 193 8. 938 1. 00 13. 63 A 46. 626 73. 334 8. 485 1. 00 15. 44 A 45. 228 73. 903 8. 815 1. 00 14. 59 A 44. 153 72. 900 8. 383 1. 00 12. 94 A 45. 110 74. 183 10. 304 1. 00 15. 69 A 46. 875 74. 188 6. 975 1. 00 16. 91 A 46. 681 71. 96	C C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1978 1979 1980 1981 1982 1983 1984 1985 1986 1987 1988 1990 1991 1992 1993	CB CG1 CG2 C O N CA CB CG OD1 ND2 C O N CA	VAL VAL VAL VAL VAL ASN ASN ASN ASN ASN ASN THR THR THR	271 271 271 271 271 272 272 272 272 272	46. 726 71. 746 5. 067 1. 00 16. 54 A 47. 928 70. 879 4. 646 1. 00 19. 07 A 47. 911 69. 548 5. 400 1. 00 20. 07 A 47. 878 70. 635 3. 131 1. 00 18. 62 A 45. 456 71. 041 4. 641 1. 00 15. 09 A 44. 912 70. 226 5. 383 1. 00 13. 46 A 44. 988 71. 394 3. 449 1. 00 15. 17 A 43. 812 70. 802 2. 832 1. 00 14. 94 A 43. 231 71. 767 1. 797 1. 00 13. 83 A 42. 010 71. 205 1. 093 1. 00 14. 46 A 41. 822 69. 989 1. 007 1. 00 16. 67 A 41. 175 72. 090 0. 581 1. 00 15. 74 A 44. 310 69. 542 2. 110 1. 00 15. 93 A 44. 755 69. 617 0. 967 1. 00 16. 88	C C C C C O N C C O N C C C O C C O C C O C C O C C O C C O C C O C C O C C C O C C C O C C C C O C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	1995 (1996 (1997 (1998 1999 (1998	CG2 CON CA CB CCB CCG CDD1 CON CA	THR THR ASP ASP ASP ASP ASP	273 273 273 274 274 274 274 274 274 274 274 275 275	43. 201 65. 794 3. 471 1. 00 19. 69 A 45. 481 66. 083 4. 266 1. 00 19. 20 A 44. 009 66. 870 0. 813 1. 00 19. 92 A 44. 550 66. 154 -0. 028 1. 00 21. 20 A 42. 811 67. 424 0. 634 1. 00 20. 50 A 42. 032 67. 193 -0. 584 1. 00 20. 30 A 40. 578 67. 629 -0. 390 1. 00 21. 02 A 39. 705 66. 529 0. 178 1. 00 23. 48 A 38. 543 66. 823 0. 527 1. 00 26. 38 A 40. 168 65. 375 0. 275 1. 00 23. 88 A 42. 573 67. 870 -1. 832 1. 00 19. 89 A 42. 131 67. 556 -2. 932 1. 00 22. 08 A 43. 508 68. 802 -1. 676 1. 00 18. 13 A 44. 073 69. 490 -2. 834 1. 00 19. 37 A	O C C C C O N C C C C C C C C C C C C C

					FIG. 4-42		(Continued)
ATOM	2009	0G	SER	275	45. 197 71. 121 -1. 444 1. 00 24. 82	·	0
ATOM	2010	C	SER	275	45. 397 68. 885 -3. 314 1. 00 19. 53	A	C
ATOM	2011	ŏ	SER	275	45. 883 69. 226 -4. 394 1. 00 19. 59	A	Ö
ATOM	2012	Ň	LEU	276	45. 971 67. 986 -2. 516 1. 00 19. 83	A	N N
ATOM	2013	CA	LEU	276	47. 241 67. 348 -2. 846 1. 00 20. 72	A	
ATOM	2014	CB	LEU	276	47. 545 66. 226 -1. 849 1. 00 19. 96	A	C C C C
ATOM	2015	CG	LEU	276	47. 725 66. 641 -0. 392 1. 00 20. 47	A	Č
ATOM	2016		LEU	276	47. 991 65. 410 0. 456 1. 00 21. 68	A	C
ATOM	2017		LEU	276	48. 875 67. 622 -0. 277 1. 00 18. 56	A	Č
ATOM	2018	C	LEU	276	47. 360 66. 790 -4. 263 1. 00 22. 34	A	č
ATOM	2019	ŏ	LEU	276	48. 290 67. 137 -4. 994 1. 00 24. 63	A	Õ
ATOM	2020	Ň	SER	277	46. 434 65. 925 -4. 656 1. 00 22. 80	A	N N
ATOM	2021	CA	SER	277	46. 501 65. 325 -5. 983 1. 00 23. 82	A	C
ATOM	2022	CB	SER	277	45. 456 64. 219 -6. 121 1. 00 22. 59	A	č
ATOM	2023	ŌĞ	SER	277	44.148 64.756 -6.044 1.00 23.44	A	ŏ
ATOM	2024	Č	SER	277	46. 305 66. 341 -7. 097 1. 00 24. 47	A	č
ATOM	2025	0	SER	277	46. 699 66. 104 -8. 231 1. 00 26. 86	A	ŏ
ATOM	2026	N	SER	278	45. 698 67. 472 -6. 768 1. 00 25. 44	A	Ň
ATOM	2027	CA	SER	278	45. 431 68. 522 -7. 745 1. 00 26. 20	A	Ċ
ATOM	2028	CB	SER	278	44.051 69.121 -7.471 1.00 25.70	A	Č
ATOM	2029	OG	SER	278	43. 831 70. 266 -8. 266 1. 00 30. 53	Ä	Ö
ATOM	2030	C	SER	278	46. 495 69. 630 -7. 739 1. 00 25. 70	A	Č ·
ATOM	2031	0	SER	278	46.603 70.414 -8.683 1.00 23.48	Ä	ő
ATOM	2032	N	VAL	279	47. 277 69. 692 -6. 672 1. 00 26. 01	Ä	Ň
ATOM	2033	CA	VAL	279	48. 327 70. 696 -6. 565 1. 00 28. 42	Ä	Ĉ
ATOM	2034	CB	VAL	279	48. 073 71. 634 -5. 350 1. 00 29. 96	A	č
ATOM	2035	CG1	VAL	279	49. 372 72. 211 -4. 834 1. 00 32. 19	A	č
ATOM	2036	CG2	VAL	279	47. 148 72. 768 -5. 776 1. 00 29. 00	Ā	Č ·
ATOM	2037	C	VAL	279	49.704 70.043 -6.470 1.00 28.21	Ä	Č
ATOM	2038	0	VAL	279	49.834 68.872 -6.088 1.00 29.00	Ā	0
ATOM	2039	Ν.	THR	280	50.728 70.801 -6.848 1.00 26.67	A	N
ATOM	2040	CA	THR	280	52. 092 70. 306 -6. 832 1. 00 26. 53	Α	C
ATOM	2041	CB	THR	280	53. 023 71. 217 -7. 645 1. 00 27. 22	Α	Č
ATOM	2042		THR	280	52. 533 71. 331 -8. 986 1. 00 29. 98	Α	0
ATOM	2043		THR	280	54. 422 70. 645 -7. 674 1. 00 26. 85	Α	C
ATOM	2044	C .	THR	280	52.618 70.254 -5.418 1.00 26.01	Α	C
ATOM	2045	0	THR	280	53. 184 69. 255 -4. 986 1. 00 27. 33	Α	0
ATOM	2046	N	ASN	281	52.402 71.341 -4.696 1.00 25.17	Α	N
ATOM	2047	CA	ASN	281	52.876 71.474 -3.334 1.00 23.78	Α	C
ATOM	2048	CB	ASN	281	54. 190 72. 250 -3. 388 1. 00 22. 28	Α	C
ATOM	2049	CG	ASN	281	54. 925 72. 287 -2. 071 1. 00 22. 87	Α	C
ATOM	2050		ASN	281	54.603 71.576 -1.116 1.00 20.83	Α	0
ATOM	2051		ASN	281	55. 948 73. 136 -2. 056 1. 00 22. 18	A	N
ATOM	2052	C	ASN	281	51. 818 72. 211 -2. 506 1. 00 23. 12	A	C
ATOM	2053	0	ASN	281	51. 876 73. 431 -2. 362 1. 00 22. 47	A	0
ATOM	2054	N	ALA	282	50. 849 71. 460 -1. 982 1. 00 23. 33	Α	N
ATOM	2055	CA	ALA	282	49. 763 72. 018 -1. 166 1. 00 23. 40	A	C
ATOM	2056	CB	ALA	282	48. 952 70. 895 -0. 547 1. 00 23. 19	A	Ç
ATOM	2057	C	ALA	282	50.320 72.912 -0.071 1.00 24.45	Α	Ċ

					TO 1	0 4	4.0			(Continued)	
					F 1	G. 4	- 43				
ATOM ATOM	2058 2059	0 N	ALA THR	282 283	51. 180 49. 817		0. 694 0. 024	1.00 25.49 1.00 24.70	A A	0 N	
ATOM ATOM	2060 2061	CA CB	THR	283	50. 326		1.021	1.00 25.33	A	C	
ATOM	2062	0G1	THR THR	283 283	50. 209 48. 834		0. 539 0. 353	1.00 27.36 1.00 29.84	A A	C 0	
ATOM	2063	· CG2		283	50.947	76. 730	-0.785	1.00 30.06	Α	С	
ATOM ATOM	2064 2065	C 0	THR THR	283 283	49. 710 48. 487		2. 406 2. 578	1.00 24.49 1.00 24.13	A	C	
ATOM	2066	N	SER	284	50. 593		3. 396	1.00 24.13	A A	O N	
ATOM	2067	CA	SER	284	50.200	74.872	4. 791	1.00 19.88	A	C	
ATOM ATOM	2068 2069	CB OG	SER SER	284	51.317		5. 624	1.00 15.88	A	C	
ATOM	2009	C	SER	$\begin{array}{c} 284 \\ 284 \end{array}$	51. 413 49. 906		5. 350 5. 288	1.00 14.23 1.00 19.24	A A	0 C	
ATOM	2071	0	SER	284	50. 774		5. 253	1.00 18.08	A	ŏ	
ATOM	2072	N	ILE	285	48. 674		5. 745	1.00 17.36	A	N	
ATOM ATOM	2073 2074	CA CB	ILE ILE	285 285	48. 249 46. 754		6. 242 5. 977	1.00 16.16 1.00 16.93	A A	C C	
ATOM	2075		ILE	285	46. 384		6. 324	1.00 10.55	A	C	
ATOM	2076		ILE	285	46. 434		4. 513	1.00 14.89	Α	C	
ATOM ATOM	2077 2078	CD1 C	ILE ILE	285 285	47. 230 48. 496	78. 526 77. 848	3. 528 7. 733	1.00 15.03 1.00 16.46	A	C	
ATOM	2079	ŏ	ILE	285	48. 116	76. 963	8. 489	1.00 10.40	A A	C 0	
ATOM	2080	N	GLN	286	49.130	78.923	8.159	1.00 16.66	A	N	
ATOM ATOM	2081 2082	CA CB	GLN GLN	286 286	49. 428 50. 778	79.088	9.563	1.00 16.43	A	C	
ATOM	2083	CG	GLN	286	51.184	79. 776 80. 070	9. 717 11. 135	1.00 16.31 1.00 17.85	A A	C C	
ATOM	2084	CD	GLN	286	52. 552	80.713	11.196	1.00 21.44	A	č	
ATOM	2085	OE1		286	53.072	81.005	12. 277	1.00 24.09	A	0	
ATOM ATOM	2086 2087	C	GLN GLN	286 286	53. 149 48. 360	80. 939 79. 885	10. 028 10. 289	1.00 19.13 1.00 16.82	A A	N C	
ATOM	2088	Ŏ	GLN	286	47. 794	80. 844	9.754	1.00 10.02	A	0	
ATOM	2089	N	ILE	287	48.070	79. 453	11.507	1.00 15.99	Α	N	
ATOM ATOM	2090 2091	CA CB	ILE ILE	287 287	47. 116 46. 036	80. 137 79. 182	12. 355 12. 894	1.00 15.11 1.00 14.14	A	C	
ATOM	2092		ILE	287	45. 147	79. 916	13. 875	1.00 14.14	A A	C C	
ATOM	2093		ILE	287	45. 206	78. 621	11.742	1.00 13.29	Α	C	
ATOM ATOM	2094 2095	CDI	ILE ILE	287 287	44. 111 47. 991	77. 675 80. 625	12. 202 13. 506	1.00 14.31 1.00 15.35	A	C	
ATOM	2096	Ö	ILE	287	48. 349	79, 860	14. 401	1.00 13.33	A A	C 0	
ATOM	2097	N	THR	288	48. 367	81.894	13.452	1.00 15.01	Ä	N	
ATOM ATOM	2098 2099	CA CB	THR THR	288 288	49. 215 49. 688	82. 465	14.482	1.00 16.71	A	C	
ATOM	2100	OG1	THR	288	48. 548	83. 874 84. 679	14. 093 13. 779	1.00 17.36 1.00 21.17	A A	C 0	
ATOM	2101	CG2	THR	288	50.621	83. 813	12.881	1.00 17.64	Ä	С	
ATOM ATOM	2102 2103	C 0	THR THR	288 288	48.510	82. 553	15.818	1.00 16.02	A	C	
ATOM	2103	N	ALA	289	47. 287 49. 301	82. 668 82. 488	15.888 16.881	1.00 16.28 1.00 16.31	A A	O N	
ATOM	2105	CA	ALA	289	48.787	82.582	18. 232	1.00 16.67	A	C	
ATOM	2106	CB	ALA	289	49.887	82. 262	19. 207	1.00 18.89	Α	C	

					FIG	. 4	- 44			(Continued)
ATOM	2107	С	ALA	289	48. 280	34. 001	18. 467	1.00 18.0		C
ATOM	2108	0	ALA	289		34. 927	17. 733	1.00 19.1		0
ATOM	2109	N	PRO	290		34. 193	19.487	1.00 18.60		N
ATOM	2110	CD	PRO	290		33. 189	20. 388	1.00 18.3		C
ATOM	2111	CA	PRO	290		35. 526	19.783	1.00 19.04		C
ATOM	2112	CB	PRO	290		85. 234	20.777	1.00 17.5		C .
ATOM	2113	CG	PRO	290		34. 055	21.499	1.00 19.73		C
ATOM	2114	C	PRO	290		86. 447	20. 369	1.00 20.4		C
ATOM	2115	0	PRO	290		85. 995	21.092	1.00 22.14		0
ATOM	2116	N	ALA	291		37. 735	20.054	1.00 19.8		N C
ATOM	2117		ALA	291		88. 728	20. 543	1.00 19.2		C
ATOM	2118	CB	ALA	291		90.132	20. 213	1.00 17.3		C
ATOM	2119	C	ALA	291		88. 610		1.00 19.6		C
ATOM	2120	0 N	ALA	291		88. 791	22. 489	1.00 21.5		0 N
ATOM	2121	N	SER	292		88. 305	22. 825 24. 264	1.00 19.10 1.00 19.90		N C
ATOM	2122	CA	SER	292		88. 185 87. 983	24. 204	1.00 19.9		C C
ATOM	2123	CB	SER	292		86.839	24. 487	1.00 19.9		
ATOM	2124 2125	OG C	SER SER	292 292		87. 055	24. 618	1.00 24.3		C C
ATOM	2126	0	SER	292		86.948	25. 760	1.00 20.2		0
ATOM ATOM	2127	N	MET	292		86. 214	23. 635	1.00 20.0		N N
ATOM	2128	CA	MET	293 293		85. 104	23. 818	1.00 20.00		Č
ATOM	2129	CB	MET	293		83. 830	23. 149	1.00 13.1		Ċ
ATOM	2130	CG	MET	293		33. 168	23. 797	1.00 15.9		Ċ
ATOM	2131	SD	MET	293		32. 503	25. 424	1. 00 15. 8		Š
ATOM	2132	CE	MET	293		82. 993	26. 296	1.00 16.4		Č .
ATOM	2133	C	MET	293		35. 487	23. 161	1.00 20.2		č
ATOM	2134	ŏ	MET	293		85. 221	23. 693	1.00 21.1		ŏ
ATOM	2135	N	LEU	294		86. 116	21.995	1.00 20.4		Ň
ATOM	2136	ĊA	LEU	294		86. 532	21. 255	1.00 21.3		Ĉ
ATOM	2137	CB	LEU	294		87. 104	19.900	1.00 21.1		č
ATOM	2138	CG	LEU	294		86.092	18.944	1.00 23.6		č
ATOM	2139		LEU	294		86.820	17.747	1. 00 22. 6		č
ATOM	2140		LEU	294		85.064	18. 493	1.00 20.9		Č
ATOM	2141	C	LEU	294		87. 533	21.981	1.00 22.0		Č
ATOM	2142	Ŏ	LEU	294	54. 953 `		21.564	1.00 23.3		0
ATOM	2143	N	ILE	295		88.156	23.053	1.00 21.8		N
ATOM	2144	CA	ILE	295		89.122	23.792	1.00 22.2		C
ATOM	2145	CB	ILE	295		89. 938	24.835	1.00 24.9		C
ATOM	2146		ILE	295		90.536	24.196	1.00 25.0		C
ATOM	2147		ILE	295		89.034	25.998	1.00 25.5		C
ATOM	2148		ILE	295		89.761	27.085	1.00 26.4		C
ATOM	2149	C	ILE	295		88.426	24.565	1.00 21.9		C
ATOM	2150	0	ILE	295		89.064	25.006	1.00 23.9		0
ATOM	2151	N	GLY	296		87.119	24. 749	1.00 20.6	5 A	N
ATOM	2152	CA	GLY	296	56.174	86.401	25.482	1.00 18.9		C
ATOM	2153	C	GLY	296		84.922	25. 167	1.00 18.4		С
ATOM	2154	0	GLY	296		84. 503	24. 202	1.00 18.6		0
ATOM	2155	N	ASP	297	56.878	84. 132	25.967	1.00 16.5	8 A	N

					FI	G. 4	- 45			(Cor	ntinued)
ATOM ATOM ATOM ATOM ATOM ATOM	2156 2157 2158 2159 2160 2161	0D2 C	ASP ASP ASP ASP ASP	297 297 297 297 297 297	56. 918 57. 960 59. 366 59. 553 60. 284 55. 553	82. 694 82. 032 82. 378 82. 882 82. 134 82. 096	25. 751 26. 650 26. 253 25. 128 27. 063 26. 041	1.00 16.95 1.00 18.00 1.00 18.62 1.00 18.23 1.00 21.29 1.00 16.02	A A A A A	C C C O C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2162 2163 2164 2165 2166 2167 2168 2169	ND1	ASP HIS HIS HIS HIS HIS	297 298 298 298 298 298 298 298	54. 847 55. 190 53. 901 52. 846 53. 245 52. 921 54. 127 54. 327	82. 537 81. 079 80. 449 81. 207 81. 448 80. 793 82. 442 82. 392	26. 942 25. 279 25. 460 24. 661 23. 241 22. 099 22. 876 21. 572	1. 00 16. 36 1. 00 14. 79 1. 00 16. 82 1. 00 14. 81 1. 00 15. 31 1. 00 14. 85 1. 00 13. 01 1. 00 14. 39	A A A A A	0 N C C C C N	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2170 2171 2172 2173 2174 2175 2176		HIS HIS HIS TYR TYR TYR TYR	298 298 298 299 299 299 299	53. 608 53. 956 55. 008 52. 802 52. 675 52. 666 53. 811	81. 400 79. 008 78. 519 78. 348 76. 963 76. 029 76. 176	21. 076 24. 979 24. 560 25. 031 24. 609 25. 816 26. 790	1.00 14.38 1.00 17.54 1.00 15.53 1.00 17.25 1.00 16.58 1.00 15.77 1.00 17.03	A A A A A A	C N C O N C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2177 2178 2179 2180 2181 2182 2183	CD1 CE1 CD2	TYR TYR TYR TYR TYR TYR TYR	299 299 299 299 299 299 299	55. 095 56. 119 53. 586 54. 600 55. 865 56. 863 51. 351	75. 762 75. 807 76. 653 76. 700 76. 270 76. 261 76. 741	26. 456 27. 380 28. 081 29. 009 28. 656 29. 595 23. 893	1. 00 14. 29 1. 00 15. 79 1. 00 15. 17 1. 00 15. 67 1. 00 15. 90 1. 00 16. 73 1. 00 17. 76	A A A A A A	C C C C C C C C C C C C C C C C C C C	·
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2184 2185 2186 2187 2188 2189 2190	O N CA CB	TYR LEU LEU LEU LEU LEU	299 300 300 300 300 300 300	50. 349 51. 355 50. 130 50. 413 49. 232 48. 131 49. 692	77. 411 75. 799 75. 413 74. 923 74. 296 75. 322	24. 178 22. 959 22. 292 20. 878 20. 139 19. 972	1.00 16.87 1.00 16.20 1.00 16.36 1.00 16.40 1.00 14.78 1.00 16.55 1.00 15.08	A A A A A A	O N C C C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2191 2192 2193 2194 2195 2196	C O N CA CB SG	LEU LEU CYS CYS CYS CYS	300 300 301 301 301 301 301	49. 777 50. 568 48. 629 48. 288 48. 208 46. 943 47. 032	74. 243 73. 312 74. 290 73. 202 73. 722 74. 962 72. 399	23. 205 23. 335 23. 873 24. 782 26. 220 26. 503 24. 468	1. 00 17. 58 1. 00 17. 21 1. 00 19. 46 1. 00 22. 20 1. 00 22. 63 1. 00 26. 56 1. 00 23. 29	A A A A A A	C O N C C S	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2198 2199 2200 2201 2202 2203	O N CA CB	CYS ASP ASP ASP ASP ASP	301 302 302 302 302 302 302	46.690	71. 481 72. 731 71. 976 72. 223 71. 355 70. 170 71. 844	25. 210 23. 386 23. 015 23. 991 23. 680 24. 066 23. 029	1. 00 25. 66 1. 00 23. 55 1. 00 24. 19 1. 00 26. 49 1. 00 28. 68 1. 00 30. 65 1. 00 30. 37	A A A A A	0 N C C C	V

(Continued)

F	T	G.	4 -	- 4	6
1.		u.	-		•

ATOM	2205	C	ASP	302	44.658	72.292	21.610	1.0023.22	A	C
	2206		ASP	302	44. 523	73.455	21. 226	1.00 24.26	Α	0
ATOM						71. 237	20.857	1.00 21.65	Ä	Ň
ATOM	2207		VAL	303	44. 385					
ATOM	2208	CA	VAL	303	43.902	71.349	19. 493	1.00 20.79	A	C
ATOM	2209	CB	VAL	303	44.926	70.803	18.480	1.00 21.88	A	C
ATOM	2210		VAL	303	44. 420	71.028	17.051	1.00 20.34	Α	С
					46. 273	71.465	18. 702	1.00 20.12	Ā	C
ATOM	2211		VAL	303						
ATOM	2212	C	VAL	303	42.657	70.494	19.417	1.00 20.38	Ą	C
ATOM	2213	0	VAL	303	42.687	69.306	19. 744	1.00 19.45	A	0
ATOM	2214	N	THR	304	41.562	71.102	18.982	1.00 20.04	A	N
ATOM	2215	CA	THR	304	40. 302	70.394	18.882	1.00 19.30	A	С
ATOM	2216	CB	THR	304	39. 494	70.546	20. 191	1.00 19:73	A	C
						70.024	21. 287	1.00 20.19	Ä	ŏ
ATOM	2217	0G1	THR	304	40. 256					
ATOM	2218	CG2	THR	304	38. 168	69.812	20.090	1.00 17.51	A	C
ATOM	2219	С	THR	304	39. 467	70.930	17. 733	1.00 18.56	A	C
ATOM	2220	0	THR	304	39. 185	72.127	17.674	1.00 19.32	A	0
ATOM	2221	Ň	TRP	305	39.082	70.042	16.819	1.00 18.08	A	N
ATOM	2222	CA	TRP	305	38. 243	70.422	15.681	1.00 16.88	Ä	Ċ
								1.00 13.92		č
ATOM	2223	CB	TRP	305	38. 332	69.394	14. 546		A	
ATOM	2224	CG	TRP	305	39. 581	69.464	13.745	1.00 13.82	A	C
ATOM	2225	CD2	TRP	305	39. 815	70.296	12.606	1.00 13.04	A	C
ATOM	2226	CE2	TRP	305	41.143	70.068	12.189	1.00 13.12	A	C
ATOM	2227	CE3	TRP	305	39. 031	71.216	11.899	1.00 13.55	A	C
		CD1	TRP	305	40. 745	68. 781	13. 967	1.00 13.51	Ä	č
ATOM	2228								_	
ATOM	2229	NE1	TRP	305	41.688	69. 138	13.036	1.00 11.41	A	N
ATOM	2230	CZ2	TRP	305	41.704	70.729	11.094	1.00 12.03	A	C
ATOM	2231	CZ3	TRP	305	39.591	71.873	10.809	1.00 14.16	A	C
ATOM	2232	CH2	TRP	305	40.914	71.625	10.419	1.00 13.92	A	C
ATOM	2233	C	TRP	305	36. 803	70.477	16. 155	1.00 16.35	A	Ċ
		_	TRP	305	36. 368	69. 613	16.917	1.00 16.55	Ä	ŏ
ATOM	2234	0								
ATOM	2235	N	ALA	306	36.064	71.484	15.704	1.00 16.10	A	N
ATOM	2236	CA	ALA	306	34. 661	71.620	16.079	1.00 17.20	A	C
ATOM	2237	CB .	ALA	306	34.336	73.074	16.384	1.00 18.47	A	C
ATOM	2238	C	ALA.	306	33.770	71.110	14.956	1.00 16.79	A	C
ATOM	2239	Ŏ	ALA	306	32.829	70.369	15. 191	1.00 18.46	A	0
	2240	N	THR	307	34. 076	71.516	13. 733	1.00 18.36	A	Ň
ATOM										
ATOM	2241	CA	THR	307	33. 314	71.100	12.564	1.00 18.83	Ą	C
ATOM	2242	CB	THR	307	$32.\ 387$	72.222	12.072	1.00 18.43	A	C
ATOM	2243	0G1	THR	307	33. 178	73.254	11.473	1.00 20.76	A	0
ATOM	2244	CG2	THR	307	31.593	72.811	13.225	1.00 16.72	Α	C
ATOM	2245	C	THR	307	34. 299	70.778	11.442	1.00 20.34	Α	C
		ŏ	THR	307	35. 494	70.626	11.689	1.00 22.05	A	Õ.
ATOM	2246									
ATOM	2247	N	GLN	308	33. 798	70. 688	10.213	1.00 20.11	A	N
ATOM	2248	CA	GLN	308	34.640	70. 389	9.066	1.00 19.71	A	C
ATOM	2249	CB	GLN	308	33. 799	69.942	7.866	1.00 19.44	Α	C
ATOM	2250	CG	GLN	308	32.845	68. 791	8.118	1.00 21.53	A	C
ATOM	2251	CD	GLN	308	33. 524	67. 505	8. 557	1.00 23.81	Α	C
	2252	0E1	GLN	308	32. 854	66. 565	9.003	1.00 25.80	Ä	ŏ
ATOM										
ATOM	2253	NEZ	GLN	308	34.848	67. 449	8.430	1.00 21.04	Α	· N

				FIG. 4-47	(Continued)
ATOM	2254	C GLN	308	35.440 71.616 8.653 1.00 19.98 A	С
ATOM	2255	O GLN	308	36. 421 71. 501 7. 922 1. 00 21. 84 A	0
ATOM	2256	N GLU	309	35. 022 72. 789 9. 114 1. 00 19. 41 A	N
ATOM	2257	CA GLU	309	35.710 74.019 8.751 1.00 20.93 A	C
ATOM	2258	CB GLU	309	34. 920 74. 764 7. 685 1. 00 21. 98 A	C
ATOM	2259	CG GLU	309	34.709 73.971 6.419 1.00 26.38 A	C
ATOM	2260	CD GLU	309	33. 890 74. 731 5. 413 1. 00 29. 11 A	C
ATOM	2261	OE1 GLU	309	33. 665 74. 192 4. 305 1. 00 31. 98 A	0
ATOM	2262	OE2 GLU	309	33. 471 75. 869 5. 736 1. 00 28. 78 A	0
ATOM	2263	C GLU	309	35. 924 74. 939 9. 932 1. 00 21. 37 A	C
ATOM ATOM	2264 2265	O GLU	309	36. 075 76. 152 9. 764 1. 00 21. 97 A	0
ATOM	2266	N ARG CA ARG	310 310	35. 941 74. 360 11. 125 1. 00 20. 65 A	N C
ATOM	2267	CB ARG	310	36. 133 75. 131 12. 340 1. 00 20. 50 A 34. 779 75. 445 12. 986 1. 00 19. 87 A	C
ATOM	2268	CG ARG	310	34. 779 75. 445 12. 986 1. 00 19. 87 A 34. 888 76. 186 14. 305 1. 00 22. 38 A	C C C
ATOM	2269	CD ARG	310	33. 519 76. 630 14. 786 1. 00 21. 66 A	C
ATOM	2270	NE ARG	310	32. 952 77. 605 13. 870 1. 00 20. 43 A	N
ATOM	2271	CZ ARG	310	31. 660 77. 884 13. 785 1. 00 19. 88 A	Č
ATOM	2272	NH1 ARG	310	30. 794 77. 261 14. 569 1. 00 21. 42 A	Ň
ATOM	2273	NH2 ARG	310	31. 235 78. 776 12. 902 1. 00 21. 69 A	N
ATOM	2274	C ARG	310	37. 009 74. 346 13. 304 1. 00 19. 05 A	Ċ
ATOM	2275	O ARG	310	36. 701 73. 214 13. 671 1. 00 20. 19 A	0
ATOM	2276	N ILE	311	38. 108 74. 959 13. 710 1. 00 17. 88 A	N
ATOM	2277	CA ILE	311	39. 044 74. 320 14. 619 1. 00 17. 41 A	С
ATOM	2278	CB ILE	311	40. 371 73. 991 13. 859 1. 00 17. 28 A	C
ATOM	2279	CG2 ILE	311	40. 982 75. 252 13. 305 1. 00 14. 23 A	C
ATOM ATOM	2280 2281	CG1 ILE	311 311	41. 358 73. 254 14. 765 1. 00 17. 79 A	C
ATOM	2282	CDI ILE	311	42. 589 72. 763 14. 011 1. 00 15. 43 A 39. 283 75. 258 15. 802 1. 00 17. 03 A	C
ATOM	2283	0 ILE	311		C
ATOM	2284	N SER	312	39. 267 76. 481 15. 649 1. 00 17. 06 A 39. 461 74. 692 16. 988 1. 00 16. 94 A	O N
ATOM	2285	CA SER	312	39. 694 75. 517 18. 163 1. 00 18. 32 A	C
ATOM	2286	CB SER	312	38. 631 75. 244 19. 235 1. 00 19. 09 A	Č
ATOM	2287	OG SER	312	39. 008 74. 173 20. 074 1. 00 18. 57 A	Ŏ
ATOM	2288	C SER	312		Č
ATOM	2289	0 SER	312	41.552 74.131 18.795 1.00 17.71 A	Ö
ATOM	2290	N LEU	313	41. 738 76. 349 19. 148 1. 00 19. 07 A	N
ATOM	2291	CA LEU	313	43. 080 76. 271 19. 708 1. 00 20. 08 A	С
ATOM	2292	CB LEU	313	44. 093 76. 931 18. 768 1. 00 19. 12 A	C .
ATOM	2293	CG LEU	313	44. 239 76. 409 17. 341 1. 00 20. 02 A	C
ATOM	2294	CD1 LEU	313	45. 480 77. 038 16. 712 1. 00 19. 82 A	Č
ATOM ATOM	2295 2296	CD2 LEU	313	44. 361 74. 892 17. 351 1. 00 20. 74 A	C
ATOM	2290 2297	C LEU O LEU	313	43. 172 76. 957 21. 062 1. 00 21. 08 A	C
ATOM	2298	O LEU N GLN	313 314	42.608 78.030 21.265 1.00 21.22 A	0
ATOM	2299	CA GLN	314	43. 898 76. 333 21. 981 1. 00 22. 23 A 44. 096 76. 884 23. 308 1. 00 22. 40 A	N C
ATOM	2300	CB GLN	314	44. 096 76. 884 23. 308 1. 00 22. 40 A 43. 545 75. 935 24. 365 1. 00 24. 62 A	C C
ATOM	2301	CG GLN	314	42. 033 75. 860 24. 406 1. 00 27. 30 A	C
ATOM	2302	CD GLN	314	41. 536 74. 832 25. 401 1. 00 29. 52 A	C
				-2, 500 12, 502 251 201 1, 00 20, 02 A	v

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		N.		FIG. 4-48	(Continued)
ATOM	2303	OE1 GLN	314	41.827 74.911 26.598 1.00 29.38 A	0
ATOM	2304	NE2 GLN	314	40. 786 73. 854 24. 911 1. 00 30. 52 A	. N
ATOM	2305	C GLN	314	45. 584 77. 099 23. 532 1. 00 22. 00 A	Ĉ
ATOM	2306	0 GLN	314	46. 382 76. 176 23. 419 1. 00 22. 34 A	0
ATOM	2307	N TRP	315	45. 954 78. 333 23. 833 1. 00 21. 50 A	N
ATOM	2308	CA TRP	315	47. 343 78. 667 24. 070 1. 00 20. 70 A	С
ATOM	2309	CB TRP	315	47. 748 79. 873 23. 226 1. 00 18. 74 A	С
ATOM	2310	CG TRP	315	47. 480 79. 711 21. 746 1. 00 17. 87 A	С
ATOM	2311	CD2 TRP	315	48. 435 79. 368 20. 733 1. 00 14. 81 A	C
ATOM	2312	CE2 TRP	315	47.764 79.419 19.491 1.00 14.29 A	C
ATOM	2313	CE3 TRP	315	49. 793 79. 029 20. 753 1. 00 13. 32 A	C .
ATOM	2314	CD1 TRP	315	46. 299 79. 936 21. 095 1. 00 15. 84 A	C
ATOM	2315	NE1 TRP	315	46. 463 79. 769 19. 742 1. 00 13. 87 A	N
ATOM	- 2316	CZ2 TRP	315	48. 407 79. 147 18. 278 1. 00 12. 51 A	С
ATOM	2317	CZ3 TRP	315	50. 433 78. 760 19. 545 1. 00 13. 87 A	C
ATOM	2318	CH2 TRP	315	49. 736 78. 822 18. 325 1. 00 12. 57 A	C
ATOM	2319	C TRP	315	47. 530 78. 976 25. 545 1. 00 21. 60 A	С
ATOM	2320	0 TRP	315	46. 615 79. 463 26. 205 1. 00 22. 41 A	0
ATOM	2321	N LEU	316	48. 721 78. 689 26. 056 1. 00 21. 81 A	N
ATOM	2322	CA LEU	316	49.033 78.915 27.458 1.00 22.64 A	C
ATOM	2323	CB LEU	316	49. 034 77. 573 28. 192 1. 00 22. 20 A	C
ATOM	$\begin{array}{c} 2324 \\ 2325 \end{array}$	CG LEU	316	49. 655 77. 484 29. 584 1. 00 23. 04 A	C
ATOM ATOM	2326	CD1 LEU CD2 LEU	316 316	48. 953 78. 438 30. 530 1. 00 24. 08 A 49. 557 76. 049 30. 085 1. 00 19. 71 A	C
ATOM	2327	CDZ LEU	316		C C
ATOM	2328	0 LEU	316		0
ATOM .	2329	N ARG	317	51. 392 79. 192 27. 046 1. 00 26. 77 A 50. 388 80. 704 28. 383 1. 00 23. 92 A	N
ATOM	2330	CA ARG	317	51.603 81.475 28.630 1.00 22.55 A	C
ATOM	2331	CB ARG	317	51. 265 82. 787 29. 337 1. 00 25. 72 A	Č
ATOM	2332	CG ARG	317	50. 490 83. 785 28. 504 1. 00 26. 56 A	č
ATOM	2333	CD ARG	317	50.187 85.012 29.327 1.00 26.99 A	č
ATOM	2334	NE ARG	317	49. 796 86. 141 28. 494 1. 00 30. 37 A	Ň
ATOM	2335	CZ ARG	317	49. 278 87. 269 28. 966 1. 00 30. 55 A	Ĉ
ATOM	2336	NH1 ARG	317	49. 082 87. 414 30. 273 1. 00 29. 99 A	Ň
ATOM	2337	NH2 ARG	317	48. 972 88. 256 28. 132 1. 00 28. 53 A	N
ATOM	2338	C ARG	317	52. 580 80, 705 29. 500 1. 00 21. 07 A	C
ATOM	2339	0 ARG	317	52. 175 79. 920 30. 359 1. 00 19. 79 A	0
ATOM	2340	N ARG	318	53. 871 80. 941 29. 290 1. 00 19. 43 A	N
ATOM	2341	CA ARG	318	54. 876 80. 259 30. 084 1. 00 17. 08 A	C
ATOM	2342	CB ARG	318	56. 263 80. 850 29. 845 1. 00 15. 15 A	C
ATOM	2343	CG ARG	318	57. 345 80. 075 30. 564 1. 00 13. 58 A	C
ATOM	2344	CD ARG	318	58. 671 80. 165 29. 853 1. 00 13. 59 A	C
ATOM	2345	NE ARG	318	59. 687 79. 341 30. 504 1. 00 11. 13 A	N
ATOM	2346	CZ ARG	318	60. 895 79. 135 30. 001 1. 00 10. 46 A	C
ATOM.	2347	NH1 ARG	318	61. 220 79. 694 28. 850 1. 00 11. 29 A	N
ATOM	2348	NH2 ARG	318	61. 773 78. 378 30. 642 1. 00 10. 86 A	N
ATOM	2349	C ARG	318	54.500 80.354 31.555 1.00 16.61 A	C
ATOM	2350	O ARG	318	54. 794 79. 448 32. 318 1. 00 20. 33 A	0 N
ATOM	2351	N ILE	319	53. 869 81. 455 31. 954 1. 00 16. 59 A	N

				FΙ	G. 4	- 49			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2355 CC 2356 CI 2357 C 2358 O 2359 N 2360 CA 2361 CI 2362 CC 2363 CI 2364 OI 2365 NI 2366 C 2367 O 2368 N 2369 CA 2370 CI 2371 CC 2371 CC 2373 NI 2374 C 2375 O 2376 N 2377 CA 2377 CA 2377 CA	B ILE G2 ILE G1 ILE G1 ILE G1 ILE GLN GLN GCN GCN GCN GCN GCN GCN GCN GCN GCN GC	319 319 319 319 319 319 320 320 320 320 320 321 321 321 321 321 321 321 321 321 321	53. 396 53. 389 52. 720 54. 828 55. 712 51. 972 51. 012 51. 870 50. 623 50. 939 52. 000 52. 304 51. 431 53. 554 49. 368 48. 645 49. 079 47. 871 48. 226 48. 776 49. 166 48. 801 46. 983 46. 095 47. 222 46. 482 47. 105	81. 607 83. 078 83. 210 83. 589 82. 743 81. 065 81. 808 79. 747 79. 001 77. 516 77. 044 75. 577 74. 734 75. 261 79. 351 78. 466 80. 633 81. 010 81. 785 83. 166 83. 491 83. 975 81. 843 82. 555 81. 715 82. 466 83. 856	33. 330 33. 776 35. 128 33. 878 34. 787 33. 251 33. 246 33. 420 32. 444 32. 570 32. 403 34. 038 34. 472 34. 207 34. 931 36. 203 35. 925 34. 804 36. 980 34. 479 32. 719 31. 719 31. 599	1. 00 17. 40 1. 00 17. 03 1. 00 17. 19 1. 00 19. 57 1. 00 19. 56 1. 00 17. 56 1. 00 18. 71 1. 00 16. 12 1. 00 14. 59 1. 00 12. 17 1. 00 10. 79 1. 00 12. 70 1. 00 13. 71 1. 00 16. 32 1. 00 14. 51 1. 00 18. 37 1. 00 19. 38 1. 00 20. 21 1. 00 23. 59 1. 00 22. 35 1. 00 27. 82 1. 00 18. 69 1. 00 19. 10 1. 00 17. 65 1. 00 18. 28 1. 00 18. 09	A A A A A A A A A A A A A A A A A A A	C C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2379 CC 2380 CI 2381 CE 2382 CI 2383 CE 2384 CZ 2385 OH 2386 C 2387 O 2388 N 2389 CA 2390 CI 2391 OC 2391 OC 2392 C 2393 O 2394 N 2395 CA 2396 CI	G TYR COLUMN TYR COLUM	322 322 322 322 322 322 322 323 323 323	46. 319 46. 561 45. 843 45. 340 44. 624 44. 163 46. 518 47. 583 45. 351 45. 237 44. 871 43. 662 44. 163 43. 250 44. 277 43. 309 43. 925 42. 944 44. 290 42. 839 43. 631	84. 856 85. 037 85. 987 85. 645 86. 589 86. 758 87. 704 81. 750 81. 587 81. 318 80. 638 79. 163 79. 025 81. 943 81. 199 81. 802 82. 995 83. 509 84. 105 80. 776 79. 985	30. 792 29. 428 28. 694 31. 401 30. 681 29. 334 28. 638 30. 363 29. 764 29. 896 28. 612 28. 806 29. 535 27. 777	1. 00 18. 09 1. 00 20. 14 1. 00 21. 33 1. 00 22. 14 1. 00 20. 00 1. 00 19. 18 1. 00 21. 74 1. 00 24. 04 1. 00 18. 70 1. 00 17. 43 1. 00 17. 45 1. 00 17. 45 1. 00 17. 51 1. 00 17. 51 1. 00 17. 88 1. 00 17. 88 1. 00 18. 20 1. 00 18. 44 1. 00 18. 83 1. 00 19. 32 1. 00 18. 46 1. 00 18. 78 1. 00 18. 78 1. 00 18. 78 1. 00 18. 75	A A A A A A A A A A A A A A A A A A A	C C C C C C C C C C C C C C C C C C C

					(Continued)
				FIG. 4-50	,
ATOM ATOM	2401 2402	N MET CA MET	325 325	41. 549 80. 772 24. 231 1. 00 17. 55 41. 046 79. 832 23. 245 1. 00 17. 68	N C
ATOM	2403	CB MET	325	39. 832 79. 062 23. 769 1. 00 19. 82 A	C
ATOM ATOM	2404 2405	CG MET SD MET	$\begin{array}{c} 325 \\ 325 \end{array}$	39. 272 78. 043 22. 774 1. 00 20. 18 A 37. 681 77. 304 23. 268 1. 00 23. 11 A	C S
ATOM	2406	CE MET	325	38. 209 75. 734 23. 896 1. 00 24. 95 A	S C
ATOM ATOM	2407 2408	C MET O MET	$\begin{array}{c} 325 \\ 325 \end{array}$	40. 641 80. 584 21. 999 1. 00 18. 03 A 39. 932 81. 583 22. 076 1. 00 16. 88 A	0 .C
ATOM	2409	N ASP	326	41. 114 80. 118 20. 852 1. 00 18. 60 A	N
ATOM	2410	CA ASP	326	40. 749 80. 738 19. 595 1. 00 20. 69 A	C
ATOM	2411	CB ASP	326	41. 988 81. 158 18. 797 1. 00 22. 43 A	C
ATOM ATOM	$\frac{2412}{2413}$	CG ASP OD1 ASP	$\frac{326}{326}$	42. 329 82. 638 18. 970 1. 00 26. 03 A 41. 511 83. 384 19. 547 1. 00 26. 48 A	C 0
ATOM	2414	OD1 ASP	326	43.415 83.063 18.518 1.00 28.75 A	ő
ATOM	2415	C ASP	326	39. 924 79. 739 18. 800 1. 00 19. 88 A	C
ATOM	2416	0 ASP	326	40. 254 78. 563 18. 729 1. 00 21. 77 A	0
ATOM ATOM	2417 2418	N ILE CA ILE	$\begin{array}{c} 327 \\ 327 \end{array}$	38. 832 80. 208 18. 223 1. 00 20. 27 A 37. 980 79. 355 17. 419 1. 00 22. 22 A	N C
ATOM	2419	CB ILE	327	36. 529 79. 393 17. 941 1. 00 20. 50 A	C
ATOM	2420	CG2 ILE	327	35. 600 78. 697 16. 985 1. 00 19. 07 A	C
ATOM	2421 2422	CG1 ILE	$\begin{array}{c} 327 \\ 327 \end{array}$	36. 483 78. 691 19. 305 1. 00 21. 51 A 35. 164 78. 766 20. 006 1. 00 20. 97 A	C C
ATOM ATOM	2423	CDI ILE	327	35. 164 78. 766 20. 006 1. 00 20. 97 A 38. 113 79. 908 16. 015 1. 00 23. 66 A	C
ATOM	2424	0 ILE	327	37. 625 80. 984 15. 716 1. 00 26. 18 A	Ö
ATOM	2425	N CYS	328	38. 804 79. 162 15. 161 1. 00 26. 09 A	N
ATOM ATOM	$\frac{2426}{2427}$	CA CYS C CYS	328 328	39. 069 79. 608 13. 805 1. 00 26. 75 A 38. 274 78. 890 12. 721 1. 00 27. 13 A	C C
ATOM	2428	0 CYS	328	38. 274 78. 890 12. 721 1. 00 27. 13 A 38. 168 77. 663 12. 705 1. 00 27. 70 A	0
ATOM	2429	CB CYS	328	40.564 79.481 13.547 1.00 27.02 A	C
ATOM	2430	SG CYS	328	41. 567 79. 984 14. 986 1. 00 28. 23 A	S
ATOM ATOM	2431 2432	N ASP CA ASP	$\frac{329}{329}$	37. 729 79. 686 11. 807 1. 00 26. 60 A 36. 913 79. 198 10. 710 1. 00 26. 21 A	N
ATOM	2433	CB ASP	329 -		- C
ATOM	2434	CG ASP	329	34.684 79.595 11.842 1.00 26.75 A	Č
ATOM	2435	OD1 ASP	329	35. 181 79. 407 12. 969 1. 00 27. 44 A	0
ATOM ATOM	2436 2437	OD2 ASP C ASP	$\frac{329}{329}$	33. 460 79. 493 11. 625 1. 00 28. 96 A 37. 613 79. 349 9. 367 1. 00 28. 54 A	0 C
ATOM	2438	0 ASP	$\frac{323}{329}$	38. 314 80. 334 9. 120 1. 00 29. 27 A	0
ATOM	2439	N TYR	330	37. 416 78. 371 8. 492 1. 00 29. 31 A	N
ATOM	2440	CA TYR	330	38. 027 78. 411 7. 173 1. 00 29. 64 A	C
ATOM ATOM	2441 2442	CB TYR CG TYR	330 330	38. 011 77. 019 6. 542 1. 00 30. 55 A 38. 597 76. 980 5. 151 1. 00 31. 78 A	C C
ATOM	2443	CD1 TYR	330	39.919 77.367 4.919 1.00 32.26 A	Č
ATOM	2444	CE1 TYR	330	40. 460 77. 341 3. 641 1. 00 32. 18 A	C
ATOM	2445	CD2 TYR	330	37. 832 76. 561 4. 066 1. 00 32. 94 A	C
ATOM ATOM	2446 2447	CE2 TYR CZ TYR	330 330	38. 364 76. 526 2. 779 1. 00 32. 62. A 39. 676 76. 920 2. 574 1. 00 33. 67 A	C C
ATOM	2448	OH TYR	330	40.193 76.914 1.299 1.00 34.33 A	0
ATOM	2449	C TYR	330	37. 314 79. 387 6. 243 1. 00 30. 14 A	C

(((Continued)
					FIG	i. 4	- 51			
ATOM	2450	0	TYR	330	36.098	79. 313	6.058	1.00 28.65	A	0
ATOM	2451	N	ASP	331		80. 308	5.666	1.00 31.49	Α	N
ATOM	2452	CA	ASP	331	37. 511	81.262	4.730	1.00 33.80	Α	. С
ATOM	2453	CB	ASP	331	38. 191	82.618	4.862	1.00 36.63	Α	C
ATOM	2454	CG	ASP	331		83.661	3.956	1.00 39.35	Α	C
ATOM	2455		ASP	331		83. 455	2.724	1.00 40.70	Α	0
ATOM	2456		ASP	331		84.684	4. 479	1.00 42.41	Α	0
ATOM	2457	C	ASP	331		80. 696	3.336	1.00 35.29	Α	C
ATOM	2458	0	ASP	331		80. 730	2.817	1.00 35.63	Α	0
ATOM	2459	N	GLU	332		80.170	2.743	1.00 36.11	A	Ŋ
ATOM.	2460	CA	GLU	332		79. 562	1.426	1.00 37.77	A	C
ATOM	2461	CB	GLU	332		78. 970	1.080	1.00 38.87	Ą	C
ATOM	2462	CG	GLU	332		78. 510	-0.354	1.00 43.60	A	C
ATOM	2463	CD	GLU	332		77. 897	-0.620	1.00 47.15	A	C
ATOM ATOM	2464 2465	0E1		332		77.771	-1.807	1.00 48.97	A	0
ATOM	2466	C	GLU GLU	$\frac{332}{332}$		77.534	0.358	1.00 48.40	A	0
ATOM	2467	Ö	GLU	332 332		80.465	0. 293	1.00 38.19 1.00 39.73	A	C
ATOM	2468	N	SER	333		79. 982 31. 764	-0.655 0.375	1.00 39.73	A	0 N
ATOM	2469	CA	SER	333		32.652	-0. 704	1.00 31.07	A A	N C
ATOM	2470	CB	SER	333		33. 858	-0.814	1.00 38.03	A	C .
ATOM	2471	0G	SER	333		34. 795	0. 223	1.00 40.60	A	0
ATOM	2472	Č	SER	333		33. 135	-0. 577	1.00 37.74	A	Č
ATOM	2473	Ŏ	SER	333		33. 838	-1.448	1.00 38.52	A	Ö
ATOM	2474	N	SER	334		32. 761	0.506	1.00 38.49	A	N
ATOM	2475	CA	SER	334		33. 163	0.708	1.00 37.49	Ä	Ċ
ATOM	2476	CB	SER	334		34. 180	1.844	1.00 38.50	Ä	č
ATOM	2477	0G	SER	334		33. 536	3.108	1.00 38.48	A	0 .
ATOM	2478	C	SER	334		31.947	1.058	1.00 35.98	A	Č
	, 2479	0	SER	334		32.029	1.148	1.00 36.41	Α	0
ATOM	2480	N	GLY	335		30.817	1.263	1.00 35.13	A	N
ATOM	2481	CA	GLY	335		79.620	1.620	1.00 35.71	A	C
ATOM	2482	C	GLY	335		79.872	2.894	1.00 35.19	A	С
ATOM	2483	0	GLY	335		9. 201	3.172	1.00 35.61	A	0
ATOM	2484	N	ARG	336		30. 855	3.666	1.00 33.99	A	N
ATOM	2485	CA	ARG	336		31. 197	4.919	1.00 33.15	A	С
ATOM	2486	CB	ARG	336		32.696	4.991	1.00 36.78	A	C
ATOM	2487	CG.	ARG	336		33. 232	3.884	1.00 42.04	A	C
ATOM	2488	CD	ARG	336		34. 374	4.416	1.00 45.76	Ą	C
ATOM	2489	NE C7	ARG	336		35. 359	5. 147	1.00 48.92	A	N
ATOM	2490	CZ NU 1	ARG	336		6. 192	6.055	1.00 50.76	A	Ç
ATOM ATOM	2491 2492		ARG ARG	336 336		6.159	6.348	1.00 52.08	A	N
ATOM	2492	C	ARG	336		7. 057	6.675	1.00 52.33	A	N
ATOM	2493 2494	0	ARG	336		0.801	6.118	1.00 30.26	A	C
ATOM	2495	N	TRP	337		0.449	5.981	1.00 29.07	A	0 N
ATOM	2496	CA	TRP	337		0. 869 0. 531	7. 294 8. 533	1.00 26.94	A	N C
ATOM	2497	CB	TRP	337		9. 403	9. 248	1. 00 24. 29 1. 00 19. 88	A A	C
ATOM	2498	CG	TRP	337		8.074	3. 246 8. <u>56</u> 1	1.00 15.00	A	C

•					FI	G. 4	- 52			(Continued)
ATOM	2499		TRP	337	41.481	77. 077	8.861	1.00 9.80	Α	C
ATOM	2500	CE2		337	41.651	76.026	7.927	1.00 9.92	Α	C
ATOM	2501		TRP	337	40.475	76. 970	9.825	1.00 7.74	Α	C
ATOM	2502	CD1	TRP	337	43.173	77.601	7.485	1.00 12.90	Α	C
ATOM	2503	NE1	TRP	337	42.688	76.369	7.099	1.00 9.82	Α	N
ATOM	2504		TRP	337	40.849	74. 885	7. 935	1.00 9.71	A	C
ATOM	2505		TRP	337	39.675	75.836	9.832	1.00 7.79	Α	C
ATOM	2506		TRP	337	39.866	74.808	8.894	1.00 10.33	Α	C
ATOM	2507	C	TRP	337	41.783	81.758	9.425	1.00 24.55	Α	C
ATOM	2508	0	TRP	337	42. 794	82.360	9.766	1.00 26.73	Α	0
ATOM	2509	N	ASN	338	40.570	82. 128	9.806	1.00 25.00	Α	N
ATOM	2510	CA	ASN	338	40.381	83. 296	10.648	1.00 26.17	A	C
ATOM	2511	CB	ASN	338	39.464	84. 300	9.949	1.00 28.44	A	C .
ATOM	2512	CG	ASN	338	40.016	84. 761	8.612	1.00 30.42	A	C
ATOM	2513		ASN	338	39. 320	84. 711	7.596	1.00 32.04	A	0
ATOM	2514		ASN	338	41.271	85. 217	8.606	1.00 28.33	A	Ŋ
ATOM	2515	Ç	ASN	338	39.810	82. 958	12.012	1.00 25.29	A	C
ATOM	2516	0	ASN	338	38.957		12.148	1.00 25.29	A	0
ATOM	2517	N	CYS	339	40. 293	83. 668	13.023	1.00 25.00	A	N
ATOM	2518	CA	CYS	339	39. 833	83. 482	14.389	1.00 24.73	A	C
ATOM	2519	C	CYS	339	39. 289	84. 829	14.888	1.00 22.42	A	C
ATOM	2520	0	CYS	339	40.051	85. 717	15. 249	1.00 21.56	A	0
ATOM	2521	CB	CYS	339	40.992	83. 014	15. 285	1.00 25.93	A	C
ATOM	2522	SG	CYS	339	42.199	81.865	14.526	1.00 29.61	A	S
ATOM	2523	N	LEU	340	37.968	84. 978	14.889	1.00 22.38	A	N
ATOM	2524	CA	LEU	340	37. 333	86. 212	15.347	1.00 20.83	A	C
ATOM	2525	CB	LEU	340	35.839	86. 185	15.069	1.00 19.89	A	C
ATOM	2526	CC	LEU	340	35. 364	86. 201	13.626	1.00 19.14	A	C
ATOM	2527		LEU	340	33.877	85.883	13.593	1.00 19.65	A	C
ATOM ATOM	2528 2529		LEU	340 340	35.647	87. 551	13.012	1.00 19.21	A	C
ATOM	2529 2530	C	LEU LEU	340 340	37. 521	86. 406	16.835	1.00 20.16	A	C
ATOM	2530 2531	O N	VAL	340 341	37. 337 37. 866	85.478	17.615	1.00 20.80	A	0
ATOM	2532	CA	VAL	341	38. 066	87. 625 87. 949	17. 225 18. 627	1.00 20.46	A	N
ATOM	2533		VAL	341	38. 536		18. 786	1.00 20.11	A	C
ATOM	2534	CG1		341	38. 972	89. 647	20. 221	1.00 21.45 1.00 22.38	A	C
ATOM	2535	CG2		341	39. 688	89.672	17.819	1.00 24.38	A A	C C
ATOM	2536	C	VAL	341	36. 770	87. 749	19.403	1.00 24.20	A	C
ATOM	2537	ŏ	VAL	341	36. 785	87. 423	20. 585	1.00 16.31	A	0
ATOM	2538	N	ALA	342	35. 644	87. 941	18. 731	1.00 17.77	A	N N
ATOM	2539	CA	ALA	342	34. 345	87. 756	19.370	1.00 19.00	A	C
ATOM	2540	CB	ALA	342	33. 228	88. 125	18.407	1.00 13.04	A	C
ATOM	2541	CD	ALA	342	34. 177	86. 302	19.829	1.00 10.09	A	C
ATOM	2542	ŏ	ALA	342	33. 245	85. 987	20. 580	1.00 13.13	A	0
ATOM	2543	Ň	ARG	343	35. 078	85. 422	19.384	1.00 16.12	A	N
ATOM	2544	ĊA	ARG	343	35.008	84. 017	19.766	1.00 16.37	A	C
ATOM	2545	CB	ARG	343	34. 962	83. 138	18.521	1.00 18.14	A	C
ATOM	2546	CG	ARG	343	33. 726	83. 390	17.687	1.00 20.31	A	č
ATOM	2547	CD	ARG	343	33. 803	82. 695	16.357	1.00 21.82	. A	č

				FIG. 4-53	(Continued)
45000					
ATOM ATOM	2548 2549	NE ARG CZ ARG	$\begin{array}{c} 343 \\ 343 \end{array}$	32. 615 82. 969 15. 561 1. 00 23. 94 A 32. 373 82. 415 14. 383 1. 00 26. 14 A	N C
ATOM	2550	NH1 ARG	343	33. 242 81. 559 13. 864 1. 00 28. 42 A	N N
ATOM	2551	NH2 ARG	343	31. 256 82. 703 13. 734 1. 00 30. 23 A	N
ATOM	2552	C ARG	343	36.164 83.603 20.650 1.00 17.09 A	C
ATOM	2553	0 ARG	343	36. 275 82. 452 21. 057 1. 00 16. 76 A	0
ATOM ATOM	2554	N GLN	344	37. 030 84. 553 20. 955 1. 00 18. 05 A	N
ATOM	2555 2556	CA GLN CB GLN	344 344	38. 175 84. 267 21. 791 1. 00 18. 90 A 39. 191 85. 385 21. 645 1. 00 18. 03 A	C
ATOM	2557	CG GLN	344	39. 191 85. 385 21. 645 1. 00 18. 03 A 40. 585 85. 012 22. 038 1. 00 17. 99 A	C
ATOM	2558	CD GLN	344	41. 571 86. 088 21. 657 1. 00 18. 02 A	Č
ATOM	2559	OE1 GLN	344	41. 711 87. 089 22. 353 1. 00 17. 71 A	Ŏ.
ATOM	2560	NE2 GLN	344	42. 246 85. 897 20. 527 1. 00 17. 42 A	N
ATOM	2561	C GLN	344	37. 708 84. 170 23. 234 1. 00 19. 61 A	C
ATOM ATOM	2562 2563	O GLN N HIS	344 345	37. 069 85. 087 23. 730 1. 00 21. 89 A 38. 013 83. 057 23. 897 1. 00 18. 47 A	0
ATOM	2564	CA HIS	345	38. 013 83. 057 23. 897 1. 00 18. 47 A 37. 624 82. 868 25. 287 1. 00 17. 92 A	N C
ATOM	2565	CB HIS	345	36. 786 81. 600 25. 453 1. 00 16. 07 A	C
ATOM	2566	CG HIS	345	35. 478 81. 641 24. 726 1. 00 15. 01 A	č
ATOM	2567	CD2 HIS	345	34. 223 81. 895 25. 164 1. 00 14. 43 A	Č
ATOM	2568	ND1 HIS	345	35. 371 81. 420 23. 369 1. 00 15. 56 A	N
ATOM ATOM	2569 2570	CE1 HIS NE2 HIS	$\begin{array}{c} 345 \\ 345 \end{array}$	34. 108 81. 535 23. 002 1. 00 12. 57 A	C
ATOM	2571	C HIS	345	33. 390 81. 823 24. 073 1. 00 14. 20 A 38. 854 82. 789 26. 172 1. 00 19. 64 A	N C
ATOM	2572	0 HIS	345	39. 839 82. 129 25. 825 1. 00 22. 18 A	0
ATOM	2573	N ILE	346	38. 790 83. 460 27. 319 1. 00 20. 11 A	N
ATOM	2574	CA ILE	346	39. 899 83. 501 28. 264 1. 00 21. 08 A	C
ATOM	2575	CB ILE	346	40. 135 84. 928 28. 760 1. 00 20. 44 A	C
ATOM ATOM	2576 2577	CG2 ILE CG1 ILE	346	41. 357 84. 972 29. 667 1. 00 20. 95 A	C
ATOM	2578	CD1 ILE	346 346	40. 338 85. 860 27. 572 1. 00 19. 87 40. 466 87. 298 27. 978 1. 00 22. 20	C
ATOM	2579	C ILE	346	40. 466 87. 298 27. 978 1. 00 22. 20 A 39. 657 82. 624 29. 482 1. 00 23. 76 A	C C
ATOM	2580	0 ILE	346	38. 535 82. 537 29. 975 1. 00 24. 67 A	Ö
ATOM	2581	N GLU	347	40.714 81.976 29.967 1.00 25.01 A	Ň
ATOM	2582	CA GLU	347	40. 601 81. 123 31. 141 1. 00 28. 30 A	C
ATOM ATOM	2583 2584	CB GLU CG GLU	347	40. 459 79. 656 30. 733 1. 00 26. 51 A	C
ATOM	2585	CO GLU	$\frac{347}{347}$	40. 089 78. 740 31. 891 1. 00 27. 38 A 40. 169 77. 268 31. 527 1. 00 29. 51 A	C
ATOM	2586	OE1 GLU	347	40. 169 77. 268 31. 527 1. 00 29. 51 A 39. 877 76. 936 30. 359 1. 00 29. 48 A	C 0
ATOM	2587	OE2 GLU	347	40. 511 76. 439 32. 405 1. 00 29. 57 A	0
ATOM	2588	C GLU	347	41. 836 81. 288 32. 021 1. 00 30. 87 A	Č.
ATOM	2589	O GLU	347	42.865 80.661 31.777 1.00 33.35 A	0
ATOM	2590 2501	N MET	348	41. 741 82. 131 33. 044 1. 00 32. 50 - A	N
ATOM ATOM	2591 2592	CA MET CB MET	348 348	42. 877 82. 347 33. 926 1. 00 34. 46 A	C
ATOM	2593	CG MET	348	43. 215 83. 843 34. 002 1. 00 37. 48 A 42. 168 84. 723 34. 661 1. 00 41. 62 A	C C
ATOM	2594	SD MET	348	42. 028 86. 340 33. 825 1. 00 48. 03 A	S
ATOM	2595	CE MET	348	43. 541 87. 158 34. 341 1. 00 46. 60 A	Č
ATOM	2596	C MET	348	42. 628 81. 784 35. 315 1. 00 33. 55 A	Ċ

					• •	•				(Continued)
					FΙ	G. 4	- 54			(00110111111111111111111111111111111111
						04 050	05 541	1 00 04 05		0
ATOM	2597	0	MET	348		81.070		1.00 34.35	A	0
ATOM	2598	N	SER	349	43. 534	82.085	36. 235	1.00 32.30	A	N
ATOM	2599	CA	SER	349		81.623	37.612	1.00 31.26	A	C
ATOM	2600	CB	SER	349	43.961	80. 197	37. 744	1.00 31.22	A	C
ATOM	2601	0G	SER	349		79.760	39.090	1.00 32.92	A	0 .
ATOM	2602	C	SER	349		82. 573	38. 474	1.00 31.16	A	C
ATOM	2603	0	SER	349		82.950	38. 113	1.00 31.25	A	0
ATOM	2604	N	THR	350		82.962	39. 611	1.00 30.83	A	N
ATOM	2605	CA	THR	350		83. 896	40.516	1.00 28.43	. A	C
ATOM	2606	CB	THR	350		84. 938	41.027	1.00 28.93	A	C
ATOM	2607		THR	350		84. 268	41.703	1.00 27.68	A	0
ATOM	2608		THR	350	42. 751	85.733	39.864	1.00 27.87	A	C
ATOM	2609	C	THR	350	44. 971	83. 198	41.714	1.00 27.14	A	C
ATOM	2610	0	THR	350	45. 781	83. 786	42. 431	1.00 27.62	A	0
ATOM	2611	N	THR	351	44.610	81.936	41.913	1.00 25.72	A	N
ATOM	2612	CA	THR	351	45. 109	81. 161	43.035	1.00 24.77	A	C
ATOM	2613	CB	THR	351	43. 945	80.536	43. 786	1.00 25.52	A	C
ATOM	2614		THR	351	43.166	79.746	42.877	1.00 24.95	A	0
ATOM	2615		THR	351	43.069	81.617	44. 385	1.00 24.61	A	C
ATOM	2616	C	THR	351	46.081	80.047	42.659	1.00 25.48	A	C
ATOM	2617	0	THR	351	46.648	79. 392	43. 535	1.00 25.57	A	0 N
ATOM	2618	N	GLY	352	46. 261	79.825	41.361	1.00 25.19	A	N
ATOM	2619	CA	GLY	352	47.170	78. 786	40.909	1.00 24.62	A	C .
ATOM	2620	C	GLY	352	47. 371	78. 797	39. 403	1.00 24.61	A	C 0
ATOM	2621	0	GLY	352	47.417	79.853	38.774	1.00 25.15	- A	N N
ATOM	2622	N	TRP	353	47. 499	77.612	38. 825 37. 390	1.00 23.36	A	C
ATOM	2623	CA	TRP TRP	353 353	47. 684	77. 470 76. 291	37. 116	1.00 21.38 1.00 17.49	A	C
ATOM	2624	CB CG	TRP	353	48. 631 48. 272	75. 023	37. 849	1.00 17.49	A A	Č
ATOM ATOM	2625 2626		TRP	353	48. 587	74. 693	39. 209	1.00 10.34	A	C
ATOM	2627		TRP	353	48. 053	73. 409	39. 462	1.00 14.04	A	Č
ATOM	2628		TRP	353	49. 270	75. 356	40. 238	1.00 14.55	A	C
ATOM	2629		TRP	353	47. 578	73. 957	37. 351	1.00 14.33	A	Ċ
ATOM.	2630		TRP	353	47. 445	72. 985	38. 311	1.00 12.84	A	N
ATOM		CZ2		353	48. 180			1.00 14.93	A	
ATOM	2632		TRP	353	49. 398	74. 719	41.480	1.00 14.33	A	č
ATOM	2633	CH2		353	48. 853	73. 436	41.700	1.00 15.07	Ä	č
ATOM	2634	C	TRP	353	46. 303	77. 236	36. 782	1.00 22.43	Ä	č
ATOM	2635	ŏ	TRP	353	45. 307	77. 292	37. 495	1.00 22.69	Ä	ŏ
ATOM	2636	Ň	VAL	354	46. 231	76. 990	35. 479	1.00 22.83	Ä	Ň
ATOM	2637	CA	VAL	354	44. 944	76. 749	34. 836	1.00 24.15	Ä	Ċ
ATOM	2638	CB	VAL	354	44. 818	77. 513	33.498	1.00 25.09	A	č
ATOM	2639		VAL	354	43.610	77. 006	32.718	1.00 24.29	A	č
ATOM	2640		VAL	354	44. 673	79.007	33. 762	1.00 24.71	A	č
ATOM	2641	C	VAL	354	44. 799	75. 264	34. 569	1.00 24.96	A	Č
ATOM	2642	ŏ	VAL	354	45. 751	74. 628	34. 127	1.00 26.10	A	ŏ
ATOM	2643	N	GLY	355	43.609	74. 722	34. 841	1.00 24.28	A	Ň
ATOM	2644	CA	GLY	355	43. 354	73. 303	34.640	1.00 22.67	Ā	Ĉ
ATOM	2645	C	GLY	355	44.040	72.457	35.696	1.00 22.77	Α	C

					FIG. 4-55	(Continued)
ATOM	2646	0	GLY	355	44. 743 72. 989 36. 548 1. 00 22. 56 A	0
ATOM	2647	N	ARG	356	43. 843 71. 145 35. 668 1. 00 23. 29 A	N
ATOM	2648	CA	ARG	356	44. 505 70. 299 36. 654 1. 00 24. 86 A	C
ATOM	2649	CB	ARG	356	43. 927 68. 886 36. 645 1. 00 24. 91 A	C
ATOM	2650	CG	ARG	356	42. 495 68. 808 37. 122 1. 00 27. 84 A	C
ATOM	2651	CD	ARG	356	41. 973 67. 391 37. 036 1. 00 31. 58 A	C
ATOM	2652	NE	ARG	356	40.518 67.340 37.149 1.00 35.53 A 39.849 67.607 38.261 1.00 37.59 A	.N C
ATOM	2653	CZ	ARG	356		N N
ATOM	2654		ARG	356 356	40.513 67.939 39.362 1.00 40.39 A 38.520 67.547 38.272 1.00 37.65 A	N N
ATOM	2655	NH2		356	45.989 70.255 36.314 1.00 25.60 A	C
ATOM ATOM	2656 2657	C 0	ARG ARG	356	46. 844 70. 508 37. 163 1. 00 28. 06 A	0
ATOM	2658	N	PHE	357	46. 285 69. 940 35. 060 1. 00 23. 61 A	N N
ATOM	2659	CA	PHE	357	47.659 69.876 34.587 1.00 21.95 A	Č
ATOM	2660	CB	PHE	357	48.029 68.442 34.205 1.00 15.99 A	č
ATOM	2661	CG	PHE	357	48. 173 67. 524 35. 380 1. 00 12. 89 A	č
ATOM	2662	CD1		357	49. 361 67. 491 36. 115 1. 00 11. 73 A	č
ATOM	2663		PHE.	357	47. 126 66. 693 35. 763 1. 00 10. 46 A	č
ATOM	2664	CE1		357	49. 507 66. 638 37. 216 1. 00 7. 55 A	Č
ATOM	2665		PHE .		47. 263 65. 838 36. 863 1. 00 11. 70 A	Č
ATOM	2666	CZ	PHE	357	48. 459 65. 811 37. 591 1. 00 6. 24 A	Ċ
ATOM	2667	С	PHE	357	47.775 70.786 33.377 1.00 23.17 A	C
ATOM	2668	0	PHE	357	48. 877 71. 196 33. 005 1. 00 26. 25 A	0
ATOM	2669	N	ARG	358	46.626 71.100 32.782 1.00 20.84 A	N
ATOM	2670	CA	ARG	358	46.541 71.972 31.615 1.00 20.05 A	C
ATOM	2671	CB	ARG	358	47. 156 71. 297 30. 396 1. 00 19. 30 A	С
ATOM	2672	CG	ARG	358	46. 496 69. 991 30. 011 1. 00 21. 15 A	С
ATOM	2673	CD	ARG	358	46. 866 69. 613 28. 598 1. 00 24. 58 A	C
ATOM	2674	NE	ARG	358	46. 293 68. 333 28. 205 1. 00 31. 68 A	N
ATOM	2675	CZ	ARG	358	46. 163 67. 924 26. 943 1. 00 34. 22 A	C
ATOM	2676		ARG	358	46. 564 68. 701 25. 939 1. 00 31. 56 A	N
ATOM	2677		ARG.	358	45. 640 66. 727 26. 687 1. 00 33. 62 A	N
ATOM	2678	C	ARG	358	45. 081 72. 315 31. 313 1. 00 20. 40 A	C
ATOM	2679	0	ARG	358	44.168 71.608 31.734 1.00 20.47 A	0
ATOM	2680				44.840 73.404 30.570 1.00 21.33 A	
ATOM	2681	CD	PRO	359	45. 785 74. 338 29. 940 1. 00 20. 09 A 43. 455 73. 772 30. 254 1. 00 21. 44 A	C
ATOM ATOM	2682 2683	CA CB	PRO PRO	359 359		C C
ATOM	2684	CG	PRO	359	43.624 74.911 29.264 1.00 20.76 A 44.907 75.539 29.713 1.00 21.86 A	C
ATOM	2685	C.	PRO	359	42.741 72.574 29.652 1.00 21.94 A	Č
ATOM	2686	Ö	PRO	359	43. 314 71. 866 28. 827 1. 00 21. 94 A	Õ
ATOM	2687	N	SER	360	41.499 72.350 30.070 1.00 22.48 A	N .
ATOM	2688	CA	SER	360	40. 723 71. 208 29. 596 1. 00 24. 26 A	Č .
ATOM	2689	CB	SER	360	39. 501 70. 986 30. 497 1. 00 25. 29 A	č
ATOM	2690	OG ·		360	38.505 71.976 30.283 1.00 27.66 A	ŏ
ATOM	2691	Č	SER	360	40. 262 71. 280 28. 140 1. 00 25. 67 A	Č
ATOM	2692	ŏ	SER	360	40.117 72.359 27.555 1.00 25.66 A	Ö
ATOM	2693	Ň	GLU	361	40.024 70.104 27.573 1.00 25.65 A	Ň
ATOM	2694	ĊA	GLU	361	39. 581 69. 972 26. 199 1. 00 27. 20 A	
			-			

				•						(0
					ास	G. 4	- 5 6			(Continued)
						J. 1				
ATOM	2695	CB	GLU	361	39.803	68.540	25.713	1.00 30.37	Α	C
ATOM '	2696	CG	GLU	361	39. 356	67.444	26.683	1.00 36.42	Α	C
ATOM	2697	CD	GLU	361	40.340	67. 226	27.839	1.00 42.80	Α	C
ATOM	2698		GLU	361	40.317	68.002	28.822	1.00 43.77	Α	0
ATOM	2699		GLU	361	41.152	66.274	27. 757	1.00 46.60	Α	0
ATOM	2700	C	GLU	361	38.112	70.324	26.052	1.00 25.88	Α	C
ATOM	2701	0	GLU	361	37. 295	69.955	26.888	1.00 27.12	Α	0 ,
ATOM	2702	N	PRO	362	37.760	71.061	24. 989	1.00 23.97	Α	N
ATOM	2703	CD	PRO	362	38.650	71.837	24.106	1.00 23.33	Α	C
ATOM	2704	CA	PR0	362	36.365	71.436	24. 767	1.00 22.45	Α	С
ATOM	2705	CB	PRO	362	36.485	72.714	23. 945	1.00 23.21	Α	C
ATOM	2706	CG	PRO	362	37.679	72.437		1.00 21.08	Α	C
ATOM	2707	C	PRO	362	35. 621	70.338	24.013	1.00 21.91	Α	C
ATOM	2708	0	PR0	362	36.216	69.582	23. 249	1.00 22.96	Α	0
ATOM	2709	N	HIS	363	34. 318	70. 259	24. 245	1.00 21.59	Α	N
ATOM	2710	CA	HIS	363	33. 459	69.280	23.596	1.00 19.88	Α	C
ATOM	2711	CB	HIS	363	32. 868	68.353	24.649	1.00 18.03	Α	С
ATOM	2712		HIS	363	33. 898	67.568	25. 398	1.00 16.56	Α	C
ATOM	2713		HIS	363	34. 638	67.880	26.489	1.00 16.19	Α	C
ATOM	2714		HIS	363	34. 292	66.303	25.019	1.00 14.56	Α	N
ATOM	2715		HIS	363	35. 227	65.869	25.843	1.00 14.60	Α	C
ATOM	2716		HIS	363	35. 457	66.808	26. 744	1.00 16.65	Α	N
ATOM	2717	C	HIS	363	32.364	70.081	22.903	1.00 20.84	Α	C
ATOM	2718	0	HIS	363	31.535	70. 709	23.564	1.00 20.84	Α	0
ATOM	2719	N	PHE	364	32. 383	70.075	21.573	1.00 19.87	Α	N
ATOM	2720	CA	PHE	364	31.416	70.832	20. 786	1.00 18.84	Α	C
ATOM	2721	CB	PHE	364	32.042	71.310	19.470	1.00 18.67	Α	C
ATOM	2722	CG	PHE	364	33.073	72.390	19.629	1.00 18.84	Α	C
ATOM	2723		PHE	364	34. 341	72.096	20.117	1.00 17.51	Α	C
ATOM	2724		PHE	364	32.776	73. 708	19. 274	1.00 16.76	Α	C
ATOM	2725		PHE	364	35. 298	7 3. 095	20. 246	1.00 16.92	Α	C
ATOM	2726		PHE	364	33.727	74.711	19.401	1.00 16.24	Α	С
ATOM	2727	CZ	PHE	364	34. 988	74. 404	19.886	1.00 16.59	Α	C
ATOM	2728	C	PHE	364	30. 172	70.046	20. 432	1.00 19.35	Α	C
ATOM	2729	0	PHE	364	30. 226	68. 831	20. 262	1.00 20.71	Α	0
ATOM	2730	N	THR	365	29.050	70. 750	20.313	1.00 18.81	Α	N
ATOM	2731	CA	THR	365	27.805	70.113	19.912	1.00 18.11	Α	С
ATOM	2732	CB	THR	365	26.600	71.017	20.161	1.00 17.38	Α	С
ATOM	2733		THR	365	26. 521	71.991	19.119	1.00 22.40	Α	0
ATOM	2734		THR	365	26. 741	71.734	21.487	1.00 13.72	Α	С
ATOM	2735	С	THR	365	28.001	69.954	18.409	1.00 17.58	Α	С
ATOM	2736	0	THR	365	28.823	70.650	17.824	1.00 16.70	Α	0
ATOM	2737	N	LEU	366	27. 250	69.058	17. 784	1.00 19.74	Α	N
ATOM	2738	CA	LEU	366	27.388	68.799	16.350	1.00 19.89	Α	С
ATOM	2739	CB	LEU	366	26.237	67. 923	15.860	1.00 19.49	Α	С
ATOM	2740		LEU	366	26. 338	67.381	14. 431	1.00 19.63	Α	C
ATOM	2741		LEU	366	27.606	66.542	14. 282	1.00 20.45	Α	C .
ATOM	2742		LEU	366	25. 112	66.539	14.128	1.00 17.80	Α	C
ATOM	2743	C	LEU	366	27.503	70.017	15.438	1.00 21.11	Α	C

					F]	[G. 4	- 57			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2744 2745 2746 2747 2748 2749 2750 2751 2752 2753 2754 2755 2756 2757 2758 2760 2761 2762 2763 2764 2765 2766 2767 2768 2767 2768 2769 2770 2771 2772	ODO C O N CA CB CG ODI ND2 C O N CA CB CG O N CA CB CO N CA CO O N CA CO O N CA CB CC O O CB CB CB CB CB CB CB CB CB CB CB CB CB	LEU ASP ASP ASP ASP ASP GLY GLY ASN	366 367 367 367 367 367 367 368 368 368 369 369 369 369 369 369 370 370 370 370 371 371	28. 269 26. 764 26. 830 25. 567 25. 458 26. 469 24. 352 28. 047 28. 274 28. 818 30. 001 29. 740 30. 678 28. 482 28. 196 26. 838 26. 797 27. 657 25. 798 28. 270 28. 185 28. 432 28. 533 27. 145 26. 523 29. 381 29. 565 29. 910 30. 735	9 69. 989 1 71. 084 2 72. 261 3 73. 114 3 73. 796 4 74. 296 7 73. 130 7 74. 122 7 72. 772 7 73. 541 7 74. 946 7 75. 324 7 6. 669 7 77. 234 7 77. 871	14. 476 15. 722 14. 867 15. 005 16. 355 17. 094 16. 669 15. 139 14. 448 16. 155 16. 480 16. 987 17. 237 17. 164 17. 647	1.00 21.26 1.00 22.95 1.00 26.09 1.00 29.82 1.00 28.76 1.00 31.88 1.00 22.76 1.00 25.46 1.00 21.02 1.00 18.54	A A A A A A A A A A A A A A A A A A A	(Continued) O N C C C O O O C O N C C C O N C C C O N C C C O N C C C O N C C C O N C C C O N C C C O N C C C O N C C C C
ATOM ATOM ATOM	2773 2774 2775		PHE PHE PHE	371 371 371	32. 194 32. 881 32. 799	73. 808 75. 062 76. 243	23. 062 23. 546 22. 818	1.00 14.83 1.00 11.31 1.00 11.07	A A A	C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2779 2780 2781 2782 2783	CE1 CE2 CZ C O N CA	PHE PHE PHE TYR TYR	371 371 371 371 371 371 372 372	33. 635 33. 465 34. 302 34. 219 30. 703 30. 362 31. 053 31. 091	75. 050 77. 409 76. 205 77. 383 73. 545 74. 495 72. 360 72. 089	24. 726 23. 256 25. 178 24. 444 25. 048 25. 752	1. 00 11. 89 1. 00 12. 04 1. 00 9. 92 1. 00 9. 76 1. 00 16. 26 1. 00 15. 15 1. 00 16. 67 1. 00 16. 84	A A A A A A A	C C C C C O N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2785 2786 2787 2788 2789 2790 2791	CE1 CD2 CE2 CZ OH	TYR	372 372 372 372 372 372 372 372 372	30. 349 28. 892 28. 470 27. 129 27. 931 26. 592 26. 193 24. 860 32. 547	70. 801 70. 879 70. 744 70. 850 71. 124 71. 235 71. 097 71. 210 71. 977	27. 271 26. 914 25. 589 25. 255 27. 901 27. 581 26. 258 25. 944 27. 367	1. 00 16. 79 1. 00 18. 47 1. 00 16. 97 1. 00 19. 91 1. 00 18. 26 1. 00 19. 23 1. 00 21. 51 1. 00 23. 32 1. 00 18. 35	A A A A A A A	C C C C C C C C C C C C C C C C C C C

	•				_ :			(Continued)
					FIG. 4-58			
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2793 2794 2795 2796 2797 2798 2799 2800 2801 2802	O N CA CB CG CD CE NZ C	TYR LYS LYS LYS LYS LYS LYS LYS LYS LYS	372 373 373 373 373 373 373 373 373	33. 388 71. 557 26. 571 1. 00 32. 845 72. 325 28. 611 1. 00 34. 224 72. 318 29. 071 1. 00 34. 907 73. 541 28. 459 1. 00 36. 302 73. 863 28. 889 1. 00 36. 658 75. 193 28. 240 1. 00 38. 048 75. 703 28. 601 1. 00 38. 103 77. 196 28. 404 1. 00 34. 277 72. 369 30. 593 1. 00	0 20. 30 0 18. 89 0 19. 69 0 19. 69 0 20. 48 0 23. 59 0 25. 15 0 24. 26 0 20. 26 0 21. 08	A A A A A A A	O N C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2803 2804 2805 2806 2807 2808 2809 2810 2811 2812	N CA CB CG2 CG1	ILE ILE ILE ILE ILE ILE ILE ILE ILE	374 374 374 374 374 374 374 375 375	35. 215 71. 634 31. 176 1. 00 35. 358 71. 624 32. 621 1. 00 35. 960 70. 309 33. 123 1. 00 36. 100 70. 361 34. 650 1. 00 35. 095 69. 128 32. 667 1. 00 35. 652 67. 753 33. 079 1. 00 36. 290 72. 745 33. 046 1. 00 37. 408 72. 846 32. 551 1. 00 35. 824 73. 595 33. 951 1. 00	0 20. 43 0 19. 63 0 19. 72 0 19. 46 0 19. 17 0 15. 57 0 19. 75 0 21. 23 0 20. 12	A A A A A A	N C C C C C C O N
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2813 2814 2815 2816 2817 2818 2819 2820	CB CG2 CG1 CD1 C O N CA	ILE ILE ILE ILE ILE ILE ILE ILE SER SER	375 375 375 375 375 375 376 376	36. 396 76. 014 33. 700 1. 00 36. 685 75. 837 32. 215 1. 00 34. 966 76. 488 33. 919 1. 00 34. 645 77. 772 33. 186 1. 00 36. 346 74. 893 35. 929 1. 00 35. 283 74. 512 36. 426 1. 00 37. 301 75. 481 36. 634 1. 00 37. 132 75. 740 38. 051 1. 00	0 20. 15 0 20. 38 0 20. 24 0 20. 36 0 21. 00 0 21. 63 0 21. 72 0 22. 04 0 23. 67	A A A A A A A	C C C C C O N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2821 2822 2823 2824 2825 2826 2827 2828 2829		SER SER SER ASN ASN ASN ASN	376 376 376 377 377 377 377	38. 336 76. 411 40. 022 1. 00 36. 063 76. 809 38. 210 1. 00 36. 042 77. 768 37. 445 1. 00 35. 164 76. 659 39. 177 1. 00 34. 128 77. 673 39. 356 1. 00 32. 755 77. 023 39. 602 1. 00 32. 682 76. 222 40. 894 1. 00 33. 560 76. 294 41. 750 1. 00	21. 76 26. 97 24. 46 27. 59 25. 41 26. 19 25. 06 22. 15	A A A A A A A	C O C O N C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2839 2840	C C C C C C C C C C C C C C C C C C C		377 377 378 378 378 378 378 378 378 378	34. 447 78. 685 40. 456 1. 00 35. 574 78. 733 40. 960 1. 00 33. 461 79. 498 40. 822 1. 00 33. 659 80. 518 41. 845 1. 00 32. 401 81. 390 41. 988 1. 00 32. 300 82. 505 40. 939 1. 00 31. 099 83. 430 41. 148 1. 00 29. 946 82. 970 40. 972 1. 00 31. 312 84. 619 41. 489 1. 00	20. 01 28. 48 29. 51 30. 42 33. 25 36. 97 44. 33 49. 20 51. 65 50. 97 32. 75	A A A A A A A	N C O N C C C C C C

					FΙ	G. 4	- 59			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2843 2844 2845 2846 2847 2848 2849 2850 2851 2852 2853 2854 2855 2856 2857 2858 2860 2861 2862 2863 2864 2865 2866 2867 2868 2869 2870 2871 2872 2873 2874 2875 2875	OE2 CCONCABCCCCI CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	ARG	379 379 379 379 379 379 379 379 380 380 381 381 381 381 381 381 381 381 381 381	33. 842 34. 192 33. 083 31. 752 30. 678 30. 363 30. 159 35. 466 35. 952 35. 986 37. 203 36. 979 37. 935 35. 726 35. 434 34. 175 34. 394 34. 175 34. 202 34. 480 34. 955 35. 261 34. 542 35. 266 35. 855 37. 057 38. 322 39. 606 40. 647 41. 178 40. 783 42. 052	78. 687 78. 070 77. 141 77. 788 76. 751 75. 976 76. 700 77. 252 76. 712 77. 136 76. 377 74. 931 74. 167 72. 671 72. 448 71. 225 71. 035 73. 486 73. 312 72. 082 71. 909 73. 100 73. 911 72. 147 72. 003 71. 217 72. 083 73. 132 73. 449 73. 907	43. 436 44. 709 45. 182 45. 416 45. 677 44. 741 46. 815 44. 589 45. 578 43. 373 43. 171 42. 781 42. 662 42. 191 42. 903 44. 379 44. 853 46. 204 45. 296 46. 647 47. 097 48. 429 40. 678 40. 091 40. 045 38. 600 38. 081 38. 110 38. 141 38. 712 38. 096 36. 868 38. 738	1. 00 31. 75 1. 00 31. 73 1. 00 35. 37 1. 00 40. 59 1. 00 46. 30 1. 00 48. 81 1. 00 49. 11 1. 00 30. 70 1. 00 30. 56 1. 00 29. 06 1. 00 27. 19 1. 00 27. 69 1. 00 26. 46 1. 00 26. 78 1. 00 26. 62 1. 00 26. 62 1. 00 24. 99 1. 00 24. 93 1. 00 26. 88 1. 00 28. 31 1. 00 26. 88 1. 00 28. 31 1. 00 28. 94 1. 00 24. 10 1. 00 24. 10 1. 00 24. 10 1. 00 23. 35 1. 00 23. 31 1. 00 22. 46 1. 00 22. 46 1. 00 22. 46 1. 00 22. 46	A A A A A A A A A A A A A A A A A A A	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	2876 (2877 (2878 N 2879 (2880 (2881 (2882 (2883 N 2884 (2885 N 2886 (2887 (2888 N 2889 (2889 (2888 N 2889 (2889 (2888 N 2889 (2888 N 2888 N 2888 N 2888 (2888 N 2888 N 2888 N 2888 (2888 N 2888 N 2888 N 2888 N 2888 (2888 N 2888 N 2888 N 2888 N 2888 (2888 N 2888 N 2888 N 2888 N 2888 (2888 N 2888 N 2888 N 2888 N 2888 (2888 N 2888 N 2888 N 2888 N 2888 (2888 N 2888	CONTRACTOR OF THE CONTRACTOR O	ARG ARG HIS HIS HIS HIS HIS HIS HIS	382 382 383 383 383 383 383 383 383 383	34. 548 34. 189 33. 840 32. 545 31. 440 31. 177 31. 590 30. 418 30. 374 31. 076 32. 404	73. 907 71. 359 70. 270 72. 068 71. 647 72. 370 71. 797 72. 189 70. 661 70. 380 71. 291 71. 930 72. 608 71. 420 71. 589 70. 419	38. 738 38. 186 38. 645 37. 313 36. 813 37. 581 38. 939 40. 168 39. 132 40. 422 41. 073 35. 330 34. 728 34. 748 33. 329 32. 802	1. 00 22. 46 1. 00 20. 92 1. 00 18. 12 1. 00 20. 45 1. 00 20. 33 1. 00 20. 76 1. 00 22. 34 1. 00 21. 75 1. 00 20. 42 1. 00 22. 25 1. 00 20. 36 1. 00 19. 84 1. 00 19. 26 1. 00 17. 93 1. 00 17. 52	A A A A A A A A A A A A A	N C O N C C C C N C N C O N C C C

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				FI	G. 4	- 6 0			(Continued)
ATOM ATOM	2892 (CG2 ILE	384 384	30. 005 30. 928	70.566 69.097	31. 290 33. 155	1.00 15.28 1.00 12.97	A A	C C
ATOM		CD1 ILE	384	30. 093	67.865	32.909	1.00 9.57	Α	С
ATOM		C ILE	384	30. 376	72.898	33.028	1.00 19.30	Α	С
ATOM) ILE	384	29. 333	73.198	33.605	1.00 18.50	Α	0
ATOM		V CYS	385	30. 950		32. 120	1.00 21.14	A	Ŋ
ATOM		CA CYS	385	30. 349	74. 953	31. 745	1.00 24.26	A	<u>C</u> .
ATOM		C CYS	385	29. 932	74. 887	30. 284	1.00 23.62	A	C
ATOM		CYS	385	30.654	74. 334	29. 464	1.00 23.61	A	0
ATOM		CB CYS	385	31.344	76.106	31. 958	1.00 27.85	A	Ç .
ATOM ATOM		SG CYS	385	30. 561	77.640	32. 569	1.00 37.75	A	S
ATOM		N TYR CA TYR	386 386	28. 760	75. 440	29. 973	1.00 23.26	A	N
ATOM		CB TYR	386	28. 237 26. 726	75. 470	28. 609	1.00 21.88	A	C
ATOM		G TYR	386	26. 120	75. 271 75. 183	28.612	1.00 21.89	A	C
ATOM		D1 TYR	386	24. 912	75. 825	27. 228 26. 930	1.00 23.48 1.00 23.55	A	C
ATOM		E1 TYR	386	24. 323	75. 712	25.665	1.00 23.33	A	C
ATOM		D2 TYR	386	26. 728	74. 424	26. 223	1.00 24.11	A A	C C
ATOM		E2 TYR	386	26. 144	74. 299	24. 956	1.00 22.10	A	C
ATOM		Z TYR	386	24. 943	74.946	24. 686	1.00 24.39	A	C
ATOM		H TYR	386	24. 358	74. 823	23. 449	1.00 23.13	A	ő
ATOM	2912 · C		386	28. 549	76. 816	27. 962	1.00 22.02	A	č
ATOM	2913 0		386	28. 187	77.868	28. 493	1.00 22.52	Ä	ŏ
ATOM	2914 N	PHE	387	29. 201	76.775	26.806	1.00 21.19	A	Ň
ATOM	2915 C	A PHE	387	29. 582	77.988	26.080	1.00 19.95	Ä	Ĉ
ATOM		B PHE	387	31. 087	77.987	25.781	1.00 17.05	A	C
ATOM		G PHE	387	31.970	78.222	26.973	1.00 14.01	A	Č
ATOM		D1 PHE	387	32. 547	79.469	27. 185	1.00 9.81	A	С
ATOM		D2 PHE	387	32. 293	77. 178	27.835	1.00 11.20	A	С
ATOM		E1 PHE	387	33. 440	79.672	28. 231	1.00 9.80	A	С
ATOM		E2 PHE	387	33. 185	77. 376	28.885	1.00 10.91	A	C C C C
ATOM		Z PHE	387	33. 762	78. 626	29.082	1.00 9.32	Α	С
ATOM	2923 C		387	28. 888	78. 153	24. 727	1.00 20.94	A	C
ATOM ATOM	2924 0 2925 N		387	28. 552	77. 180	24.055	1.00 19.77	A	0
ATOM		GLN A GLN	388 388	28. 706		24. 332	1.00 21.79	A	N
ATOM		B GLN	388	28. 151 27. 024	79. 742	23. 030	1.00 22.21	Α.	C
ATOM		G GLN	388	25. 745	80. 760 80. 343	23. 177 22. 477	1.00 23.86	A	C
ATOM		D GLN	388	25. 096	79. 126	23. 109	1.00 29.81 1.00 32.86	A	C
ATOM		E1 GLN	388	24. 357	78. 391	22. 452	1.00 34.80	A A	C
ATOM		E2 GLN	388	25. 356	78. 913	24. 395	1.00 34.36	A	O N
ATOM	2932 C		388	29. 403	80. 382	22. 427	1.00 21.72	A	C
ATOM	2933 0		388	29. 845	81. 428	22. 893	1.00 21.72	A	0
ATOM	2934 N		389	29. 982	79. 745	21.415	1.00 20.66	A	N N
ATOM	2935 C		389	31. 231	80. 215	20. 821	1.00 21.00	Ä	C
ATOM	2936 C	B ILE	389	31.466	79. 617	19. 422	1.00 20.76	Ä	Č
ATOM		G2 ILE	389	31.410	78. 100	19.496	1.00 19.50	Ä	č
ATOM		G1 ILE	389	30.448	80.165	18.429	1.00 19.48	Ä	č
ATOM	2939 C	D1 ILE	389	30.813	79.864	16.992	1.00 19.12	A	č
									•

(Continued) FIG. 4-61 **ATOM** 2940 C ILE 389 31.483 81.713 20.735 1.00 23.29 \mathbf{c} A 82.146 **ATOM** 2941 0 ILE 32.640 20.776 389. 1.00 22.48 0 Α **ATOM** 2942 N **ASP** 390 30.423 82.505 20.611 1.00 24.96 A N 30.584 ATOM 2943 CA ASP 390 83.953 20.533 1.00 26.49 C A 2944 29.932 1.00 29.09 ATOM CB ASP 390 84.508 19.275 A \mathbf{C} ATOM **ASP** 2945 CG 390 28.467 84.215 19.216 \mathbb{C} 1.00 30.91 A **ATOM** 2946 OD1 ASP 390 27.754 84.955 18.517 1.00 35.45 0 A **ATOM** 2947 OD2 ASP 28.029 83.236 390 19.858 1.00 33.49 0 A C **ATOM** 2948 **ASP** 390 30.005 84.676 21.738 1.00 26.43 C A **ATOM** 2949 0 **ASP** 29.402 390 85.735 21.603 1.00 26.54 0 A 2950 ATOM N LYS 391 30.163 22.910 84.078 1.00 27.05 A N **ATOM** 2951 CA LYS 29.707 391 84.679 24.150 1.00 28.81 C A ATOM 2952 CB LYS 391 28.348 84.128 24.566 1.00 28.62 C **ATOM** 2953 CG LYS 27. 203 391 84.790 23.824 1.00 31.00 C A 2954 **ATOM** CD LYS 391 25.867 84.228 24.256 1.00 34.06 C Α 2955 **ATOM** CE LYS 24.733 84.772 391 23.413 1.00 33.69 \mathbb{C} A ATOM 2956 NZ LYS 1.00 36.51 391 23.454 84.073 23.742 N A ATOM C 30.772 2957 LYS 391 84.369 25.183 1.00 29.11 A C 2958 ATOM 0 LYS 391 31.192 83.223 25.327 1.00 29.45 0 A **ATOM** 2959 N LYS 31.219 85.401 392 25.888 1.00 29.66 A N **ATOM** 2960 CA LYS 392 32.281 85.248 26.872 1.00 30.67 Α **ATOM** 2961 CB LYS 392 33.069 86.558 26. 985 1.00 28.28 C Α **ATOM** 2962 CG LYS 392 33.516 C 87.119 25.636 1.00 27.07 Α **ATOM** 2963 LYS CD 392 34.330 86.098 24.852 1.00 27.55 C A **ATOM** 2964 CE LYS 392 34.643 86.588 23.449 1.00 26.02 C Α **ATOM** 2965 NZ LYS 392 35.369 87.872 23.495 1.00 25.63 Α N **ATOM** 2966 LYS C 392 31.824 84.797 28.248 1.00 31.24 C Α **ATOM** 2967 0 LYS 392 32.637 84.679 29.162 1.00 32.17 A 0 **ATOM** 2968 N ASP 393 30.531 84.548 28.403 1.00 31.57 A N **ATOM** 2969 CA ASP 393 30.015 84.098 29.690 1.00 33.64 Ċ A **ATOM** 2970 CB ASP 393 29.052 85.134 30.271 1.00 36.88 Č A **ATOM** 2971 CG ASP 393 29.734 86.450 30.567 1.00 41.66 C A **ATOM** 2972 OD1 ASP 393 30.607 86.475 31.467 1.00 43.84 A 0 2973 **ATOM** OD2 ASP 393 29.409 87.455 29.895 1.00 44.39 0 A **ATOM** 2974 C **ASP** 393 29.309 82.761 29.546 1.00 32.46 A C 28. 294 82.666 **ATOM** 2975 0 **ASP** 393 28.859 1.00 32.91 0 **ATOM** 2976 N CYS 394 29.841 81.731 30.198 1.00 30.05 N Α **ATOM** 2977 CA CYS 394 29. 243 80.410 30.115 1.00 28.94 ${\tt C}$ Α **ATOM** 2978 **CYS** 394 28.312 80.116 31.282 1.00 27.56 A CYS **ATOM** 2979 0 394 28.262 80.858 32.258 1.00 27.11 0 Α 2980 CB **ATOM** CYS 394 30.336 79.338 30.033 1.00 31.03 A C SG **ATOM** 2981 CYS 394 31.401 79.166 31.504 1.00 34.42 . S Α **ATOM** 2982 N THR 395 27.570 79.023 31.167 1.00 25.71 N Α **ATOM** 2983 CA THR 395 26.645 78.608 32.204 1.00 25.01 C Α **ATOM** 2984 CB THR 395 25.208 78.512 31.647 1.00 25.50 C A 2985 OG1 THR ATOM 395 24.709 79.833 31.407 1.00 28.36 0 A 2986 CG2 THR **ATOM** 24.289 395 77.779 32.620 1.00 21.52 C A **ATOM** 2987 C THR 395 27.048 77. 251 32.772 1.00 24.22 Ċ Α ATOM 2988 0 THR 395 76.280 27.196 32.036 1.00 24.44 0

			٠.				•			(Continued)
			•		FΙ	G. 4	- 62			(Continueu)
ATOM	2989	N	PHE	396	27. 231	77. 185	34. 084	1.00 23.09		NT .
ATOM	2990				27. 594				A	N
ATOM	2991				28. 138	76. 182			A	C
ATOM	2992		PHE		29. 581	76. 617		1.00 22.19	A	C
ATOM	2993		1 PHE		30. 604		36. 131 35. 876		A	C
ATOM	2994		2 PHE		29. 924	77. 935	26 415		A	C
ATOM	2995		1 PHE		31. 949	76.086	36. 415 35. 908	1.00 20.97 1.00 20.26	A	C
ATOM	2996		2 PHE		31. 343	78. 331	36. 447		A	C
ATOM	2997				32 279	77. 400		1.00 21.70	. A A	C C
ATOM	2998		PHE		31. 267 32. 279 26. 373	75.008	34.764	1.00 20.21	A	C
ATOM	2999		PHE		25. 311	75. 412	35. 218	1.00 20.96	A	0
ATOM	3000		ILE		26. 523	73. 779	34. 279	1.00 18.88	Ä	N
ATOM	3001		ILE		25. 412	72. 842	34. 262	1.00 18.00	A	C
ATOM	3002	CB			25. 266	72. 165	32. 879	1.00 16.55	A	Č
ATOM	3003		2 ILE		25.350	73. 209	31.787	1.00 13.63	A	č
ATOM	3004		ILE		26.366	71.130	32.669	1.00 16.02	Ä	č
ATOM	3005		ILE		26.180	70. 327	31.402	1.00 17.85	A	č
ATOM	3006	C	ILE		25.527	71.770	35. 338	1.00 19.16	A	č
ATOM	3007	0	ILE	397	24. 787	70. 787	35.330	1.00 20.44	Ā	Ŏ
ATOM	3008	N	THR		26.480	71.956	36. 244	1.00 18.55	Ä	Ň
ATOM	3009	CA	THR		26.681	71.051	37.367	1.00 19.41	A	Ċ
ATOM	3010	CB	THR		27.624	69.858	37.051	1.00 19.56	Ā	Č
ATOM	3011		THR		28.978	70. 321	36.960	1.00 22.60	A	0
ATOM	3012		THR		27.221	69. 178	35.759	1.00 18.50	Α	С
ATOM	3013	C	THR		27. 343	71.899	38.424	1.00 20.24	Α	С
ATOM	3014	0	THR		27.979	72.903	38. 104	1.00 20.11	Α	0
ATOM	3015	N	LYS		27. 185	71.511	39. 681	1.00 22.48	Α	N
ATOM	3016	CA	LYS	399	27. 795	72. 258	40.772	1.00 23.72	Α	С
ATOM	3017	CB	LYS		27.111	73.618	40.941	1.00 24.42	Α	C
ATOM	3018	CG	LYS	399	25. 689	73. 583	41.462	1.00 27.65	Α	С
ATOM ATOM	3019	CD	LYS	399	25. 269	74. 996	41.856	1.00 30.77	A	C
ATOM	3020 3021	CE	LYS	399 200	23. 861	75.054	42.414	1.00 31.89	A	C
ATOM	3021	NZ C	LYS LYS	399 399	22.841	74. 747	41.377	1.00 35.03	A	N
ATOM	3023	0	LYS	399	27. 751	71.476 70.425	42.077	1.00 22.46	A	C
ATOM	3024	N	GLY	400	27. 125 28. 435			1.00 21.96	_	0
ATOM	3025	CA	GLY	400	28. 463	71.989	43.093	1.00 21.98	A	N
ATOM	3026	C	GLY	400	20. 403 29. 891	71.319	44.378	1.00 22.66	A	C
ATOM	3027	ŏ	GLY	400	30. 831	71.115	44. 839	1.00 24.94	A	C
ATOM	3028	N	THR	401	30. 064	71.449 70.566	44.118	1.00 26.10	A	0
ATOM	3029	CA	THR	401	31.400	70. 335	46.036 46.560	1.00 25.34	A	N
ATOM	3030	CB	THR	401	31.443	70. 535	48.095	1.00 26.41	A	C
ATOM	3031		THR	401	30.615	69. 567	48. 741	1.00 27.75	A	C
ATOM	3032		THR	401	30. 924	71. 927		1.00 31.37	A	0
ATOM	3033	C	THR	401	31. 923	68. 945	48. 448 46. 197	1.00 27.06 1.00 24.83	A	C
ATOM	3034	ŏ	THR	401	32. 027	68. 049	47. 036	1.00 24.83	A	C
ATOM	3035	Ň	TRP	402	32. 229	68. 790	44. 915	1.00 20.74	A	0 N
ATOM	3036	ĊA	TRP	402	32. 781	67. 569	44. 340	1.00 22.03	A A	N C
ATOM	3037	CB	TRP	402	31.741	66. 460	44. 268	1.00 16.39	A A	C
					1 11			10.00	Λ	U

					. F]	[G. 4	- 63			(C	ontinued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3039 3040 3041 3042 3043 3044 3045 3046 3047 3050 3051 3052 3053 3054 3055 3056 3057 3058 3060 3061 3062 3063 3064 3065 3066 3067 3068 3069 3070 3071 3072 3073 3074 3075 3076 3077 3078 3079 3080 3081	CE2CCD1 NCZ CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	ILE ILE ILE GLY GLY GLY ILE ILE ILE	402 402 402 402 402 402 402 402 403 403 403 403 403 403 403 404 404 404	30. 434 30. 037 28. 701 30. 679 29. 364 28. 318 27. 989 29. 972 28. 637 33. 208 32. 956 33. 831 34. 284 35. 776 36. 122 37. 433 37. 506 38. 384 34. 028 33. 891 33. 957	66. 886 66. 865 67. 320 66. 505 67. 425 66. 608 67. 064 67. 089 67. 089 67. 484 67. 805 68. 824 69. 522 70. 728 68. 824 69. 522 70. 728 68. 824 69. 522 70. 73 66. 516 65. 305 67. 073 66. 273 66. 273 66. 273 66. 273 66. 273 66. 732 64. 593 62. 123 62. 364 63. 218 64. 290	43. 709 42. 332 42. 278 41. 137 44. 409 43. 562 41. 078 39. 924 42. 191 40. 866 40. 926 41. 923 39. 716 39. 916 38. 508 37. 305 36. 165 34. 914 36. 595 36. 363 36. 363 36. 363 36. 363 36. 363 36. 363 36. 363 36. 363 36. 363 37. 305 36. 363 36. 363 37. 365 38.	2 1.00 19.16 3 1.00 20.21 7 1.00 18.78 7 1.00 17.97 1.00 20.57 1.00 18.32 1.00 19.71 1.00 18.98 1.00 18.12 1.00 17.78 1.00 19.48 1.00 20.26 1.00 21.69 1.00 24.57 1.00 19.74 1.00 20.05 1.00 18.47 1.00 17.63 1.00 18.38 1.00 17.63 1.00 18.38 1.00 17.63 1.00 18.38 1.00 17.63 1.00 18.38 1.00 20.05 1.00 18.38 1.00 20.05 1.00 18.38 1.00 20.05 1.00 18.38 1.00 20.05 1.00 18.38 1.00 17.63 1.00 16.28 1.00 16.28 1.00 16.28 1.00 17.43 1.00 17.43 1.00 17.49 1.00 16.93		A A A A A A A A A A A A A A A A A A A	
ATOM	3086 C	CA (GLU	408	33.000	61.176	28. 122	1.00 22.31	A	C	

										(Continued)
					FI	G. 4	- 64			
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3088 C 3089 C 3090 C 3091 C 3092 C 3093 C 3094 N 3095 C 3096 C 3097 C 3098 C 3101 C 3102 C 3103 C 3104 C 3105 C 3106 C 3107 N 3108 C 3107 N 3108 C 3109 C 3110 C	CONTRACTOR	GLU GLU GLU GLU GLU GLU GLU GLU GLU GLU	408 408 408 408 408 409 409 409 409 410 410 410 410 410 411 411 411 411 411	F I 32. 691 33. 457 34. 963 35. 519 35. 594 32. 262 32. 743 31. 100 30. 356 29. 797 29. 235 28. 651 28. 937 27. 911 28. 559 27. 634 26. 959 28. 434 26. 959 28. 434 26. 998 27. 453 25. 701 24. 741 23. 902 23. 017 24. 797 23. 846 23. 971 22. 952 22. 061 21. 206	59. 922 59. 860 59. 947 59. 081 60. 877 61. 097 60. 455 61. 729 61. 685 60. 294 62. 708 63. 041	- 6 4 28. 944 30. 254 30. 048 29. 337 30. 596 26. 780 25. 846 26. 671 25. 414 25. 180 25. 386 26. 413 24. 195 24. 038 23. 796 23. 617 24. 935 23. 134 22. 879 21. 758 23. 150 22. 100 22. 418 23. 498 22. 811 22. 050 22. 882 21. 074 20. 945 19. 687	1. 00 21. 64 1. 00 23. 48 1. 00 26. 15 1. 00 28. 40 1. 00 25. 87 1. 00 22. 35 1. 00 23. 83 1. 00 22. 21 1. 00 20. 74 1. 00 21. 17 1. 00 20. 05 1. 00 19. 25 1. 00 19. 28 1. 00 19. 28 1. 00 20. 83 1. 00 20. 92 1. 00 20. 83 1. 00 20. 92 1. 00 20. 84 1. 00 19. 86 1. 00 19. 86 1. 00 19. 86 1. 00 15. 82 1. 00 15. 79 1. 00 14. 12 1. 00 20. 16 1. 00 21. 79 1. 00 20. 25 1. 00 21. 79 1. 00 20. 25 1. 00 21. 09 1. 00 22. 27	A A A A A A A A A A A A A A A A A A A	C C C O O C O N C C C C C C C C O N C C O C C O N C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3117 Oct 3118 C 3119 Oct 3120 N 3121 C 3122 C 3123 C 3124 Oct 3125 Oct 3127 Oct 3128 N 3129 C 3130 C 3131 C 3132 C 3133 C 3134 C	A B G D1 A B G D1 B1 D2	SER SER SER SER ASP ASP ASP ASP ASP ASP TYR TYR TYR TYR TYR TYR TYR	412 412 412 413 413 413 413 413 413 413 414 414 414	21. 206 20. 474 21. 158 20. 598 21. 015 20. 138 19. 036 18. 161 17. 153 18. 474 20. 822 20. 306 21. 974 22. 672 22. 369 20. 925 20. 402 19. 071 20. 074 18. 740	65. 827 64. 618 66. 118 67. 185 65. 054 65. 104 64. 047 64. 243 63. 515 65. 111 64. 918 65. 363 64. 259 63. 998 62. 572 62. 822 62. 621 61. 629 61. 424	19. 687 19. 721 22. 153 22. 379 22. 934 24. 097 23. 975 22. 751 22. 635 21. 904 25. 442 26. 470 25. 444 27. 155 27. 520 28. 714 29. 052 26. 666 26. 993	1. 00 22. 27 1. 00 25. 03 1. 00 21. 84 1. 00 22. 97 1. 00 22. 56 1. 00 24. 36 1. 00 30. 28 1. 00 32. 47 1. 00 31. 81 1. 00 24. 37 1. 00 25. 08 1. 00 24. 23 1. 00 23. 61 1. 00 25. 79 1. 00 26. 31 1. 00 26. 99 1. 00 24. 67 1. 00 25. 53	A A A A A A A A A A A A A A A A A A A	C O C C C C C C C C C C C C C C C C C C

				FIG. 4-65	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3145 CI 3146 C 3147 O 3148 N 3149 CA 3150 CI 3151 CC 3152 CI	TYR TYR TYR LEU A LEU B LEU COLLEU COLLEU TYR A TYR A TYR B TYR COLLEU TYR TYR TYR TYR TYR	414 414 415 415 415 415 415 416 416 416 416	F I G. 4 - 6 5 18. 246 61. 923 28. 188 1. 00 28. 30 A 16. 925 61. 731 28. 531 1. 00 31. 69 A 24. 180 64. 174 26. 639 1. 00 22. 81 A 24. 811 64. 040 25. 582 1. 00 22. 74 A 24. 741 64. 469 27. 809 1. 00 20. 51 A 26. 174 64. 630 27. 996 1. 00 18. 28 A 26. 502 66. 079 28. 358 1. 00 16. 58 A 27. 945 66. 406 28. 745 1. 00 14. 79 A 28. 184 67. 892 28. 606 1. 00 13. 01 A 28. 208 65. 943 30. 163 1. 00 14. 04 A 26. 518 63. 684 29. 149 1. 00 18. 57 A 25. 926 63. 763 30. 230 1. 00 18. 31 A 27. 449 62. 769 28. 909 1. 00 19. 11 A 27. 843 61. 796 29. 924 1. 00 19. 69 A 27. 963 60. 407 29. 309 1. 00 18. 66 A 26. 698 59. 926 28. 645 1. 00 17. 78 A 26. 297 60. 438 27. 410 1. 00 16. 67	C O C O N C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3153 CE 3154 CI 3155 CE 3156 CZ 3157 OH 3158 C 3159 O 3160 N 3161 CA 3162 CE 3163 CG 3164 CD 3165 CE 3166 CD	21 TYR 22 TYR 22 TYR 23 TYR 24 TYR 25 TYR 26 TYR 26 TYR 27 TYR 27 TYR 27 TYR 27 TYR 27 TYR	416 416 416 416 416 416 417 417 417 417 417 417	25. 137 59. 971 26. 786 1. 00 18. 58 A 25. 908 58. 944 29. 245 1. 00 16. 22 A 24. 754 58. 475 28. 636 1. 00 16. 35 A 24. 374 58. 986 27. 406 1. 00 18. 54 A 23. 252 58. 489 26. 784 1. 00 19. 53 A 29. 167 62. 178 30. 540 1. 00 20. 71 A 30. 117 62. 499 29. 822 1. 00 22. 92 A 29. 238 62. 138 31. 866 1. 00 19. 27 A 30. 472 62. 506 32. 544 1. 00 19. 08 A 30. 408 63. 981 32. 970 1. 00 18. 38 A 29. 383 64. 282 34. 049 1. 00 17. 93 A 29. 721 64. 213 35. 399 1. 00 15. 25 A 28. 784 64. 476 36. 391 1. 00 13. 14 A 28. 071 64. 622 33. 718 1. 00 17. 72	C C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3168 CZ 3169 OH 3170 C 3171 O 3172 N 3173 CA 3174 CB 3175 CG 3176 CG	TYR TYR TYR ILE ILE	417 417 417 417 418 418 418 418 418 418 419 419 419 419	27. 120 64. 885 34. 710 1. 00 15. 27 A 27. 488 64. 808 36. 040 1. 00 14. 25 A 26. 556 65. 046 37. 020 1. 00 14. 06 A 30. 768 61. 615 33. 747 1. 00 18. 77 A 29. 918 60. 853 34. 207 1. 00 18. 74 A 31. 996 61. 706 34. 236 1. 00 17. 63 A 32. 429 60. 926 35. 379 1. 00 16. 60 A 33. 626 60. 019 35. 015 1. 00 15. 54 A 34. 482 59. 737 36. 241 1. 00 14. 33 A 33. 107 58. 729 34. 378 1. 00 15. 75 A 34. 183 57. 767 33. 964 1. 00 15. 48 A 32. 827 61. 909 36. 453 1. 00 18. 54 A 33. 535 62. 875 36. 190 1. 00 20. 83 A 32. 670 62. 556 38. 764 1. 00 2	C C C C C C C C C C C C C C C C C C C

										(Continued)
					FΙ	G. 4	- 66			
ATOM	3185	0	SER	419	32. 783	60. 503	39. 988	1.00 20.32	Α	0
ATOM	3186	N	ASN	420	33. 152	62.427	41.107	1.00 19.64	Α	N
ATOM	3187	CA .	ASN	420	33.357	61.786	42.387	1.00 20.07	Α	C
MOTA	3188	CB	ASN	420	34. 773	62.053	42.863	1.00 18.49	A	C
ATOM	3189	CG	ASN	420	35.099	63. 518	42.872	1.00 20.69	A	C
ATOM	3190		ASN	420	34. 210	64. 358	42. 741	1.00 21.49	A	0
ATOM	3191		ASN	420	36. 376	63. 844	43.034	1.00 21.39	A	N
ATOM	3192	C	ASN	420	32. 350	62.368	43. 379	1.00 20.90	A	C
ATOM	3193	0 N	ASN	420	32.677	62.610	44. 535	1.00 21.17	A	0 N
ATOM	3194	N	GLU	421	31.127	62.600 63.160	42.914	1.00 21.68 1.00 24.26	A	N C
ATOM ATOM	3195 3196	CA CB	GLU GLU	421 421	30. 081 28. 935	63. 722	43. 761 42. 901	1.00 24.20	A A	C
ATOM	3197	CG	GLU	421	27.714	64. 214	43. 701	1.00 25.32	A	Č
ATOM	3198	CD	GLU	421	26.604	64.817	42.824	1.00 26.02	A	Č
ATOM	3199		GLU	421	25. 563	65. 237	43. 373	1.00 24.11	Ä.	ŏ
ATOM	3200		GLU	421	26. 762	64. 873	41.588	1.00 27.22	Ä	ŏ
ATOM	3201	Č	GLU	421	29.512	62. 133	44. 729	1.00 24.93	Ä	Č
ATOM	3202	0	GLU	421	29.185	62.457	45.868	1.00 27.30	A	0
ATOM	3203	N	TYR	422	29.409	60.892	44. 272	1.00 23.63	Α	N
ATOM	3204	CA	TYR	422	28.837	59.826	45.075	1.00 23.67	Α	C
ATOM	3205	CB	TYR	422	28.942	58.503	44.311	1.00 23.61	Α	
ATOM	3206	CG	TYR	422	28.015	57. 415	44.813	1.00 24.39	Α	C
ATOM	3207		TYR	422	26.642	57. 637	44.936	1.00 23.87	Α	С
ATOM	3208	CE1		422	25. 781	56.618	45.347	1.00 22.11	A	C C C C C
ATOM	3209		TYR	422	28. 505	56. 147	45.120	1.00 24.53	A	C
ATOM	3210		TYR	422	27.654	55. 124	45.533	1.00 23.32	A	C
ATOM	3211	CZ	TYR	422	26.300	55. 367	45.641	1.00 23.52	A	C
ATOM	3212	OH	TYR	422	25.471	54.349	46.031	1.00 24.33	A	0
ATOM ATOM	3213 3214	C 0	TYR TYR	$\begin{array}{c} 422 \\ 422 \end{array}$	29.399	59.679	46.493	1.00 23.57	A	C
ATOM	3214 3215	N	LYS	422	30. 599 28. 492	59. 478 59. 784	46.704 47.461	1.00 23.17 1.00 23.07	A	0 N
ATOM	3216	CA	LYS	423	28. 813	59. 661	48. 878	1.00 23.07	A A	N C
ATOM	3217	CB	LYS	423	29.156	58. 205	49. 205	1.00 24.22	A	C
ATOM	3218	CG	LYS	423	27. 967	57. 266	49.009	1.00 25.11	A	Č
ATOM	3219	CD	LYS	423	28. 303	55.809	49. 276	1.00 26.55	Ä	č
ATOM	3220	CE	LYS	423	27.079	54.930	49.002	1.00 28.11	Ä	č
ATOM	3221	NZ	LYS	423	27. 302	53. 498	49.336	1.00 27.79	Ä	N
ATOM	3222	C	LYS	423	29.923	60.583	49.347	1.00 21.46	Ā	Ċ
ATOM	3223	0	LYS	423	30. 533	60.340	50.385	1.00 20.97	A	0
ATOM	3224	N	GLY	424	30.167	61.647	48.583	1.00 21.39	Α	N
ATOM	3225	CA	GLY	424	31.201	62.608	48.930	1.00 21.20	Α	C
ATOM	3226	C	GLY	424	32.606	62.034	48.961	1.00 21.98	A	C
ATOM	3227	0	GLY	424	33. 463	62. 534	49.687	1.00 22.19	A	0
ATOM	3228	N	MET	425	32. 848	60. 991	48.173	1.00 22.44	A	N
ATOM	3229	CA	MET	425	34. 161	60. 350	48.134	1.00 23.29	Ą	C
ATOM	3230	CB	MET	425	34.003	58. 826	48.056	1.00 24.14	A	C
ATOM	3231	CC	MET	425	33. 548	58. 187	49.360	1.00 25.32	A	C
ATOM	3232	SD	MET	425 425	33.092	56. 451	49.179	1.00 29.39	A	S
ATOM	3233	CE	MET	425	34. 663	55. 611	49.406	1.00 27.92	A	С

		٠			FIG.	4 - 6 7			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3276 3277 3278 3279	NH2 C O N CA CB CG OD1 ND2 C O N CA CB CG CD1 CD2 C O N CA CB CG O N CA CB CG O N CA CB CB CG O N CA CB CB CB CB CB CB CB CB CB CB CB CB CB	LEU LEU LEU TYR TYR	425 426 426 426 426 426 427 427 428 428 429 429 429 429 429 429 429 430 430 430 431 431 431 431 431 431 432 432	35. 042 60.	827 46. 986 457 45. 835 661 47. 292 215 48. 615 172 46. 262 007 47. 062 461 48. 245 019 45. 532 107 44. 343 936 46. 252 789 45. 646 838 44. 941 706 44. 642 285 44. 684 446 43. 991 970 42. 583 108 42. 289 158 41. 712 559 40. 320 167 39. 444 292 39. 108 737 40. 302 769 39. 956 145 40. 456 389 39. 756 373 40. 302 769 39. 956 145 40. 456 389 39. 756 375 39. 857 379 39. 146 388 39. 756 375 39. 857 379 39. 146 389 39. 756 375 39. 857 379 39. 146 389 39. 756 375 39. 857 379 39. 146 385 39. 622 181 40. 620 188 40. 297 186 41. 840 187 32. 230 187 32. 230 188 42. 231 189 35. 491 180 36. 172 180 36. 172 180 36. 172 181 32. 230 182 32. 230 183 35. 491 184 34. 933	1. 00 22. 06 1. 00 22. 61 1. 00 21. 75 1. 00 21. 34 1. 00 20. 07 1. 00 20. 22 1. 00 19. 61 1. 00 20. 63 1. 00 23. 99 1. 00 19. 08 1. 00 18. 03 1. 00 18. 93 1. 00 18. 22 1. 00 17. 96 1. 00 18. 15 1. 00 18. 15 1. 00 19. 00 1. 00 16. 78 1. 00 19. 00 1. 00 20. 57 1. 00 22. 65 1. 00 25. 75 1. 00 26. 83 1. 00 26. 83 1. 00 26. 83 1. 00 26. 24 1. 00 15. 74 1. 00 12. 14 1. 00 17. 29 1. 00 18. 53 1. 00 17. 41 1. 00 17. 29 1. 00 18. 53 1. 00 17. 98 1. 00 17. 98 1. 00 17. 98 1. 00 17. 98 1. 00 17. 70 1. 00 17. 37 1. 00 19. 03 1. 00 17. 62 1. 00 20. 33 1. 00 21. 23	A A A A A A A A A A A A A A A A A A A	(Continued) C O N C C C C C O N C C C C O N C C C C
ATOM 3	3281	CB CG CD1	TYR TYR TYR	432 432 432	26. 848 61. 4 27. 068 61. 0 28. 320 60. 64	42 36.014 70 37.464	1. 00 22. 85 1. 00 25. 34 1. 00 24. 87	A A A	C C C

		*•				
				FIG. 4-68		(Continued)
				r 1 G. 4 - 0 C		
ATOM	3283	CE1 TYR	432	28.519 60.305 39.20	67 1.00 24.97	A C
ATOM	3284	CD2 TYR	432	26.019 61.142 38.38		A C
ATOM	3285	CE2 TYR	432	26. 205 60. 805 39. 72		A C
ATOM	3286	CZ TYR	432	27. 454 60. 388 40. 16		A Č
ATOM	3287	OH TYR	432	27.625 60.054 41.48		A O
ATOM	3288	C TYR	432	26. 102 60. 743 33. 73		A C
ATOM	3289	0 TYR	432	26.860 60.870 32.77		A Ö
ATOM	3290	N LYS	433	24. 802 61. 022 33. 69		A N
ATOM	3291	CA LYS	433	24. 133 61. 505 32. 49	1.00 20.98	A C
ATOM	3292	CB LYS	433	23. 290 60. 386 31. 87	6 1.00 21.14	A C
ATOM	3293	CG LYS	433	22. 564 60. 827 30. 61		A C
ATOM	3294	CD LYS	433	21.843 59.701 29.90		A C
ATOM	3295	CE LYS	433	20.643 59.235 30.68		A C
ATOM	3296	NZ LYS	433	19.801 58.370 29.81		A N
ATOM	3297	C LYS	433	23. 228 62. 687 32. 83		A C
ATOM	3298	0 LYS	433	22. 367 62. 587 33. 70		A O
ATOM	3299	N ILE	434			A N
ATOM	3300	CA ILE	434	22.591 64.980 32.41		A C
ATOM	3301	CB ILE	434	23. 427 66. 225 32. 81		A Č
ATOM	3302	CG2 ILE	434	24. 412 66. 582 31. 71		A C
ATOM	3303	CG1 ILE	434	22. 491 67. 404 33. 08		A C
ATOM	3304	CD1 ILE	434	23. 171 68. 591 33. 69		A C
ATOM	3305	C ILE	434	21.782 65.297 31.17		A C
ATOM	3306	0 ILE	434	22. 274 65. 154 30. 05		A O
ATOM	3307	N GLN	435	20.538 65.716 31.37		A N
ATOM	3308	CA GLN	435	19.666 66.034 30.24		A C
ATOM	3309	CB GLN	435	18. 202 65. 851 30. 64		A C
ATOM	3310	CG GLN	435	17. 227 66. 030 29. 49		A Č
ATOM	3311	CD GLN	435	15. 802 65. 806 29. 92		A C
ATOM	3312	OE1 GLN	435	15.446 64.720 30.37		A O
ATOM	3313	NE2 GLN	435	14.978 66.839 29.81		A N
ATOM	3314	C GLN	435	19.891 67.450 29.74		A C
ATOM	3315	0 GLN	435	19.600 68.419 30.43		A O
ATOM	3316	N LEU	436	20.401 67.564 28.52		A N
ATOM	3317	CA LEU	436	20.679 68.865 27.95		A C
ATOM	3318	CB LEU	436	21.152 68.714 26.50		A C
ATOM	3319	CG LEU	436	22.456 67.939 26.33		A Č
ATOM	3320	CD1 LEU	436	22. 938 68. 116 24. 91		A C
ATOM	3321	CD2 LEU	436	23.510 68.437 27.31		A C
ATOM	3322	C LEU	436	19. 491 69. 812 28. 02		A C
ATOM	3323	0 LEU	436	19.672 71.016 28.16		A O
ATOM	3324	N SER	437	18. 280 69. 268 27. 92		A N
ATOM	3325	CA SER	437	17.059 70.075 27.97		A C
ATOM	3326	CB SER	437	15. 925 69. 340 27. 26		A C
ATOM	3327	OG SER	437	16. 241 69. 151 25. 90		A O
ATOM	3328	C SER	437	16.610 70.437 29.39		A C
ATOM	3329	0 SER	437	15.805 71.352 29.57		A O
ATOM	3330	N ASP	438	17.124 69.714 30.38		A N
ATOM	3331	CA ASP	438	16.772 69.955 31.784		A C
				222		0

			FIG. 4-69	(Continued)
ATOM	9999 CD ACT	, ,,,,		
ATOM	3332 CB ASF 3333 CG ASF		15. 468 69. 226 32. 123 1. 00 38. 49 A 14. 996 69. 498 33. 543 1. 00 41. 58 A	C .
ATOM	3334 OD1 ASP		17 000 00 115 01 100 1 00 10 07	C
ATOM	3335 OD2 ASP		10 700 00 705 00 505	0
ATOM	3336 C ASP		13. 796 69. 785 33. 725 1. 00 43. 71 A 17. 904 69. 470 32. 700 1. 00 35. 28 A	0 C
ATOM	3337 0 ASP		18.019 68.274 32.993 1.00 33.70 A	0
ATOM	3338 N TYR	439	18. 723 70. 412 33. 158 1. 00 34. 27 A	N
ATOM	3339 CA TYR		19.862 70.105 34.013 1.00 33.69 A	Č
ATOM	3340 CB TYR		20. 740 71. 343 34. 175 1. 00 32. 29 A	č
ATOM	3341 CG TYR		21. 262 71. 886 32. 867 1. 00 30. 75 A	Č
ATOM	3342 CD1 TYR		21.565 71.028 31.810 1.00 30.00 A	C C C C C
ATOM	3343 CE1 TYR		22.071 71.516 30.611 1.00 28.65 A	C
ATOM ATOM	3344 CD2 TYR		21. 480 73. 253 32. 691 1. 00 28. 95 A	С
ATOM	3345 CE2 TYR 3346 CZ TYR		21. 987 73. 749 31. 496 1. 00 27. 97 A	С
ATOM	3347 OH TYR		22. 281 72. 875 30. 462 1. 00 27. 72 A 22. 803 73. 350 29. 284 1. 00 28. 72 A	
ATOM	3348 C TYR		10 540 40 500	0
ATOM	3349 0 TYR	439	00 400 00 010	. C
ATOM	3350 N THR	440	20. 435 69. 045 36. 076 1. 00 33. 49 A 18. 285 69. 612 35. 806 1. 00 34. 13 A	O N
ATOM	3351 CA THR	440	17. 917 69. 076 37. 115 1. 00 34. 14 A	C
ATOM	3352 CB THR	440	16. 561 69. 624 37. 609 1. 00 33. 49 A	Č
ATOM	3353 OG1 THR	440	15. 507 69. 114 36. 780 1. 00 32. 29 A	ŏ
ATOM	3354 CG2 THR	440	16.559 71.144 37.571 1.00 30.29 A	č
ATOM	3355 C THR	440	17. 794 67. 572 36. 953 1. 00 33. 89 A	Č
ATOM ATOM	3356 O THR 3357 N LYS	440	17. 684 66. 829 37. 929 1. 00 35. 16 A	0
ATOM	3357 N LYS 3358 CA LYS	441	17. 808 67. 141 35. 697 1. 00 32. 21 A	N
ATOM	3359 CB LYS	441 441	17. 703 65. 735 35. 362 1. 00 30. 32 A	C
ATOM	3360 CG LYS	441	16. 871 65. 573 34. 088 1. 00 33. 16 A 15. 369 65. 490 34. 331 1. 00 36. 13 A	C
ATOM	3361 CD LYS	441	14 040 00 001 00 100	C
ATOM	3362 CE LYS	441	19 447 00 000 00 00	C
ATOM	3363 NZ LYS	441	13. 447 66. 392 35. 649 1. 00 41. 94 A 12. 953 67. 501 36. 517 1. 00 44. 46 A	C N
ATOM	3364 C LYS	441	19. 089 65. 119 35. 179 1. 00 28. 77 A	C
ATOM	3365 0 LYS	441	19. 668 65. 159 34. 088 1. 00 28. 32 A	Ŏ
ATOM	3366 N VAL	442	19. 618 64. 564 36. 263 1. 00 25. 14 A	Ň
ATOM	3367 CA VAL	442	20. 922 63. 929 36. 243 1. 00 24. 37 A	Ċ
ATOM ATOM	3368. CB VAL 3369 CG1 VAL	442	21. 960 64. 717 37. 091 1. 00 24. 82 A	C
ATOM	3369 CG1 VAL 3370 CG2 VAL	442	23. 266 63. 936 37. 178 1. 00 22. 99 A	C
ATOM	3371 C VAL	$\begin{array}{c} 442 \\ 442 \end{array}$	22. 216 66. 084 36. 469 1. 00 23. 65 A	C
ATOM	3372 0 VAL	442	20. 786 62. 525 36. 807 1. 00 24. 10 A 20. 327 62. 341 37. 931 1. 00 22. 60 A	C
ATOM	3373 N THR	443	21 100 01 500 00 011	0
ATOM	3374 CA THR	443	21 100 00 140 00 410 1 00 00	N
ATOM	3375 CB THR	443	21. 109 60. 149 36. 419 1. 00 22. 78 A 20. 352 59. 306 35. 375 1. 00 23. 02 A	C .
ATOM	3376 OG1 THR	443	19. 017 59. 802 35. 222 1. 00 27. 68 A	0
ATOM	3377 CG2 THR	443	20. 301 57. 862 35. 800 1. 00 22. 12 A	Č
ATOM	3378 C THR	443	22. 493 59. 548 36. 551 1. 00 23. 54 A	č
ATOM	3379 0 THR	443	23. 367 59. 792 35. 721 1. 00 23. 36 A	ŏ
ATOM	3380 N CYS	444	22. 701 58. 761 37. 596 1. 00 23. 18 A	N

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				FIG. 4-70	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3391 CD 3392 C 3393 O 3394 N	CYS CYS CYS LEU	444 444 444 444 445 445 445 445 445 445	23. 981 58. 104 37. 747 1. 00 24. 13 A 23. 758 56. 712 37. 157 1. 00 22. 91 A 22. 855 55. 990 37. 573 1. 00 21. 72 A 24. 396 58. 018 39. 219 1. 00 25. 50 A 26. 053 57. 282 39. 443 1. 00 30. 81 A 24. 573 56. 348 36. 175 1. 00 22. 64 A 24. 446 55. 053 35. 513 1. 00 22. 51 A 24. 799 55. 211 34. 035 1. 00 19. 29 A 24. 049 56. 349 33. 341 1. 00 19. 36 A 24. 588 56. 552 31. 934 1. 00 16. 01 A 22. 559 56. 034 33. 319 1. 00 15. 72 A 25. 308 53. 940 36. 118 1. 00 23. 32 A 25. 203 52. 783 35. 718 1. 00 24. 58 A 26. 148 54. 274 37. 087 1. 00 23. 95	C C C S N C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3395 CA 3396 CB 3397 OG 3398 C 3399 O 3400 N 3401 CA 3402 C 3403 O 3404 CB 3405 SG 3406 N 3407 CA	SER SER SER CYS	446 446 446 446 447 447 447 447 447 447	27. 028 53. 269 37. 660 1.00 23. 89 A 28. 469 53. 555 37. 222 1.00 21. 87 A 28. 882 54. 847 37. 648 1.00 20. 09 A 26. 969 53. 145 39. 175 1.00 23. 77 A 27. 361 52. 119 39. 720 1.00 24. 69 A 26. 480 54. 184 39. 845 1.00 24. 32 A 26. 382 54. 207 41. 309 1.00 26. 45 A 25. 836 52. 946 41. 997 1.00 25. 99 A 26. 441 52. 425 42. 937 1.00 24. 44 A 25. 518 55. 396 41. 763 1.00 27. 33 A 26. 225 57. 049 41. 461 1.00 34. 75 A 24. 696 52. 456 41. 528 1.00 25. 90 A 24. 056 51. 317 42. 167 1.00 24. 38 A	C C O C C C C S N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3408 CB 3409 CG 3410 CD 3411 OE 3412 OE 3413 C 3414 O 3415 N 3416 CA 3417 CB 3418 CG	GLU GLU GLU C2 GLU GLU GLU LEU LEU GLU	448 448 448 448 448 448 449 449 449 449	22. 581 51. 637 42. 334 1. 00 23. 47 A 22. 332 53. 075 42. 721 1. 00 24. 60 A 22. 848 53. 416 44. 108 1. 00 27. 44 A 22. 617 54. 562 44. 559 1. 00 29. 17 A 23. 478 52. 548 44. 751 1. 00 28. 81 A 24. 201 49. 941 41. 537 1. 00 23. 54 A 23. 722 48. 970 42. 104 1. 00 22. 25 A 24. 844 49. 844 40. 377 1. 00 23. 78 A 25. 024 48. 547 39. 717 1. 00 23. 34 A 25. 988 48. 678 38. 548 1. 00 20. 76 A 25. 680 49. 712 37. 472 1. 00 21. 20 A 26. 872 49. 807 36. 543 1. 00 20. 05 A	C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3420 CD 3421 C 3422 O 3423 N 3424 CA 3425 CE 3426 CG 3427 OD	LEU LEU LEU ASN A ASN B ASN G ASN	449 449 449 450 450 450 450 450 450	24. 424 49. 335 36. 711 1. 00 24. 61 A 25. 551 47. 456 40. 654 1. 00 24. 61 A 25. 157 46. 298 40. 549 1. 00 26. 01 A 26. 445 47. 830 41. 562 1. 00 25. 89 A 27. 040 46. 889 42. 512 1. 00 27. 02 A 27. 939 45. 913 41. 754 1. 00 27. 92 A 28. 296 44. 695 42. 572 1. 00 31. 61 A 28. 521 44. 786 43. 783 1. 00 34. 65 A 28. 363 43. 541 41. 912 1. 00 31. 27 A 27. 877 47. 731 43. 488 1. 00 26. 54 A	C C O N C C C O N

					FIG. 4-71	(Continued)
ATOM	3430	. 0	A CNI	450		
ATOM	3431		ASN PRO	450 451	29. 099 47. 637 43. 523 1. 00 26. 25 A 27. 210 48. 558 44. 303 1. 00 27. 04 A	0
ATOM	3432		PRO	451	27. 210 48. 558 44. 303 1. 00 27. 04 A 25. 762 48. 411 44. 535 1. 00 27. 72 A	N C
ATOM	3433		PRO	451	27. 796 49. 465 45. 296 1. 00 27. 49 A	C C
ATOM	3434		PRO	451	26. 579 49. 924 46. 103 1. 00 27. 21 A	Č
ATOM	3435		PRO	451	25. 638 48. 765 45. 989 1. 00 25. 73 A	č
ATOM	3436		PR0	451	28. 938 48. 983 46. 187 1. 00 28. 75 A	č
ATOM	3437		PRO	451	29.877 49.737 46.433 1.00 30.69 A	0
ATOM	3438		GLU	452	28. 873 47. 746 46. 666 1. 00 29. 54 A	N
ATOM	3439		GLU	452	29. 918 47. 228 47. 545 1. 00 30. 30 A	С
ATOM ATOM	3440 3441	CB CG	GLU	452	29. 453 45. 937 48. 232 1. 00 33. 99 A	C
ATOM	3442	CD	GLU GLU	452 452	28. 085 46. 024 48. 890 1. 00 39. 92 A	C
ATOM	3443	0E1	GLU	452 452	27. 817 44. 848 49. 813 1. 00 45. 87 A 28. 084 43. 693 49. 402 1. 00 47. 97 A	C
ATOM	3444	0E2		452	07 000 17 070 70 70	0
ATOM	3445	C	GLU	452	27. 336 45. 076 50. 948 1. 00 47. 68 A 31. 221 46. 946 46. 816 1. 00 29. 63 A	0 C
ATOM	3446	0	GLU	452	32. 308 47. 199 47. 344 1. 00 30. 27 A	0
ATOM	3447	N	ARG	453	31. 099 46. 425 45. 600 1. 00 27. 01 A	N
ATOM	3448		ARG	453	32. 244 46. 057 44. 783 1. 00 24. 90 A	Č
ATOM	3449	CB	ARG	453	31. 950 44. 728 44. 085 1. 00 23. 08 A	Č
ATOM	3450	CG	ARG	453	32. 952 44. 337 43. 018 1. 00 22. 92 A	C
ATOM	3451	CD	ARG	453	32. 602 42. 995 42. 381 1. 00 20. 49 A	C
ATOM ATOM	3452 3453	NE CZ	ARG ARG	453	33. 504 42. 688 41. 278 1. 00 18. 31 A	N
ATOM	3454		ARG	453 453	33. 439 41. 595 40. 531 1. 00 18. 93 A	C
ATOM	3455	NH2		453 453	32. 510 40. 679 40. 763 1. 00 19. 77 A 34. 302 41. 425 39. 539 1. 00 18. 87 A	N
ATOM	3456	C	ARG	453	00 005 40 004 40 000	N
ATOM	3457	Ŏ	ARG	453	32. 695 47. 071 43. 738 1. 00 25. 72 A 33. 809 46. 962 43. 222 1. 00 24. 32 A	C
ATOM	3458	N	CYS	454	31. 857 48. 054 43. 420 1. 00 25. 94 A	O N
ATOM	3459	CA	CYS	454	32. 233 49. 012 42. 385 1. 00 25. 49 A	Č
ATOM	3460	C	CYS	454	32. 038 50. 473 42. 699 1. 00 24. 24 A	č
ATOM	3461	0_	CYS	454	30. 922 50. 970 42. 688 1. 00 26. 79 A	Ö
ATOM	3462		CYS	454	31.503 48.664 41.096 1.00 26.13 A	Ċ
ATOM	3463		CYS	454	32. 156 47. 128 40. 401 1. 00 30. 12 A	S
ATOM ATOM	3464 3465		GLN	455	33. 143 51. 165 42. 942 1. 00 22. 97 A	N
ATOM	3466		GLN GLN	455 455	33. 105 52. 576 43. 276 1. 00 23. 69 A	C
ATOM	3467		GLN	455 455	33. 536 52. 761 44. 736 1. 00 23. 41 A	C
ATOM	3468		GLN	455	32. 564 52. 187 45. 761 1. 00 24. 96 A 33. 177 52. 065 47. 150 1. 00 29. 34 A	C
ATOM	3469	0E1		455	00 004 50 005 45 55	C
ATOM	3470	NE2		455	00 700 71 000 15 000	0
ATOM	3471		GLN	455	32. 790 51. 022 47. 872 1. 00 28. 59 A 33. 992 53. 425 42. 360 1. 00 24. 57 A	N C
ATOM	3472	0	GLN	455	33. 837 54. 645 42. 294 1. 00 27. 40 A	0
ATOM	3473		TYR	456	34. 919 52. 787 41. 654 1. 00 22. 57 A	N
ATOM	3474		TYR	456	35. 821 53. 510 40. 763 1. 00 21. 75 A	Ċ
ATOM	3475		TYR	456	37. 270 53. 187 41. 124 1. 00 20. 47 A	č
ATOM	3476		TYR	456	38. 267 54. 282 40. 817 1. 00 21. 27 A	C
ATOM	3477		TYR	456	38. 659 55. 193 41. 808 1. 00 20. 27 A	C
ATOM	3478	CE1	TYR	456	39.618 56.165 41.548 1.00 18.67 A	C

		(Continued)							
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3529 C 3530 C 3531 C 3532 C 3533 N 3534 C 3535 C 3536 C 3537 C 3538 C	SE I LY CA LY CB LY CG LY CD LY CE LY	R 462 R 462 R 462 S 463 S 463 S 463 S 463 S 463 S 463	30. 694 29. 494 28. 308 30. 804 30. 572 31. 153 31. 323 31. 587 33. 047 33. 972 35. 433	53. 212 52. 381 53. 145 54. 496 55. 581 54. 373 55. 536 55. 084 55. 199 54. 435 54. 724	- 7 3 20.877 20.399 20.397 20.058 20.577 18.784 17.920 16.484 16.075 17.007 16.673	1. 00 23. 70 1. 00 23. 50 1. 00 24. 06 1. 00 24. 95 1. 00 25. 95 1. 00 27. 50 1. 00 31. 80 1. 00 33. 43 1. 00 35. 54 1. 00 36. 78 1. 00 39. 20	A A A A A A A A	C C O C O N C C C C
ATOM ATOM ATOM ATOM ATOM	3540 C 3541 O 3542 N	LY	S 463 S 463 U 464	30. 226 30. 484 29. 015	54. 098 56. 602 57. 745 56. 254 57. 247	17. 641 17. 934 17. 561 18. 354 18. 410	1.00 40.26 1.00 33.39 1.00 36.36 1.00 33.23 1.00 34.54	A A A	N C O N
ATOM ATOM ATOM ATOM ATOM	3544 C 3545 C 3546 C 3547 O	B GLI G GLI D GLI E1 GLI	U 464 J 464 J 464 J 464	26. 960 27. 528 26. 578 25. 439	57. 058 57. 366 56. 961 57. 480	17. 256 15. 882 14. 772 14. 752	1. 00 39. 82 1. 00 44. 96 1. 00 48. 72 1. 00 50. 39	A A A A	C C C C
ATOM ATOM ATOM ATOM	3549 C 3550 O 3551 N 3552 C	GLI AL <i>i</i> A AL <i>i</i>	J 464 J 464 A 465 A 465	27. 186	56. 120 57. 202 57. 659 56. 636 56. 546	13. 926 19. 729 19. 814 20. 748 22. 081	1.00 50.59 1.00 32.77 1.00 32.03 1.00 31.17 1.00 29.63	A A A A	0 C O N C
ATOM ATOM ATOM ATOM ATOM	3553 Cl 3554 C 3555 O 3556 N 3557 CA	AL/ AL/ LYS A LYS	465 465 466 466	26. 015 6 25. 176 6 25. 905 6	57. 935 55. 645 55. 824 54. 678 53. 772	22. 577 22. 164 23. 042 21. 259 21. 274	1. 00 28. 36 1. 00 29. 47 1. 00 28. 66 1. 00 28. 89 1. 00 28. 97	A A A A	C C O N C
ATOM ATOM ATOM ATOM ATOM	3558 CI 3559 CC 3560 CI 3561 CE 3562 NZ	G LYS D LYS E LYS	466 466 466	24. 585 5 23. 208 5 23. 045 5 21. 632 5	53. 122 52. 509 52. 179 51. 757 60. 441	19. 899 19. 649 18. 171 17. 814 18. 404	1.00 30.98 1.00 31.77 1.00 34.52 1.00 35.82 1.00 38.42	A A A A	C C C
ATOM ATOM ATOM ATOM ATOM	3563 C 3564 O 3565 N 3566 CA 3567 CE	LYS LYS TYR TYR	466 466 467 467	24. 987 5 24. 040 5 26. 252 5 26. 599 5	2. 704 2. 126 2. 446 1. 458	22. 339 22. 869 22. 646 23. 654	1.00 28.20 1.00 27.93 1.00 26.93 1.00 26.21	A A A	N C O N C
ATOM ATOM ATOM ATOM	3568 CC 3569 CE 3570 CE 3571 CE	G TYR D1 TYR C1 TYR D2 TYR	467 467 467 467	25. 823 4 25. 550 4 24. 494 4 25. 009 4	0. 119 9. 502 9. 917 9. 373 8. 522	23. 003 22. 207 20. 903 20. 184 22. 768	1. 00 27. 94 1. 00 30. 39 1. 00 29. 93 1. 00 31. 13 1. 00 29. 73	A A A A	C C C C
ATOM ATOM ATOM ATOM ATOM	3572 CE 3573 CZ 3574 OH 3575 C 3576 O		467	23. 698 4 22. 625 4 27. 777 5	1.949	22. 060 20. 770 20. 079 24. 470 24. 064	1.00 30.29 1.00 30.97 1.00 32.01 1.00 24.00 1.00 24.63	A A A · A A	C C O C

ATOM 3577 N. TYR 468 27.969 51.370 25.641 1.00 23.06 A N ATOM 3578 CA TYR 468 29.991 51.765 26.462 1.00 22.80 A C ATOM 3579 CB TYR 468 28.801 53.043 27.249 1.00 23.81 A C ATOM 3580 CG TYR 468 27.588 53.011 28.155 1.00 24.49 A C ATOM 3581 CDI TYR 468 27.588 53.011 28.155 1.00 24.49 A C ATOM 3581 CDI TYR 468 27.588 53.011 28.155 1.00 24.49 A C ATOM 3582 CBI TYR 468 27.588 53.011 28.155 1.00 25.51 A C ATOM 3583 CD2 TYR 468 25.006 53.082 28.486 1.00 25.51 A C ATOM 3583 CD2 TYR 468 25.006 53.082 28.486 1.00 25.57 A C ATOM 3583 CD2 TYR 468 25.380 53.91 29.857 1.00 26.39 A C ATOM 3585 CZ TYR 468 25.380 53.91 29.857 1.00 25.87 A C ATOM 3585 CZ TYR 468 25.380 53.91 29.857 1.00 25.87 A C ATOM 3585 CZ TYR 468 24.304 53.334 30.695 1.00 25.95 A O ATOM 3587 C TYR 468 29.501 50.675 27.411 1.00 21.32 A C ATOM 3589 N GLN 469 30.800 50.431 27.449 1.00 22.73 A O ATOM 3589 N GLN 469 31.388 49.29 28.315 1.00 19.27 A C ATOM 3589 CA GLN 469 31.388 49.29 28.315 1.00 19.27 A C ATOM 3591 CB GLN 469 31.604 48.891 28.807 29.3815 1.00 19.27 A C ATOM 3591 CB GLN 469 33.600 79.32 28.321 1.00 21.12 A C ATOM 3595 CD GLN 469 33.600 79.28 28.315 1.00 21.12 A C ATOM 3599 CA GLN 469 33.600 79.35 28.321 1.00 21.72 A C ATOM 3599 CA GLN 469 33.600 79.79 32.801 1.00 23.85 A C ATOM 3599 CD GLN 469 33.600 79.79 32.801 1.00 23.85 A C ATOM 3599 CD GLN 469 33.600 79.79 32.801 1.00 23.85 A C ATOM 3599 CA GLN 469 33.502 24.600 27.348 1.00 23.85 A C ATOM 3599 CA GLN 469 33.502 24.600 27.348 1.00 23.85 A C ATOM 3599 CA GLN 469 33.502 24.600 27.348 1.00 23.85 A C ATOM 3599 CA GLN 469 33.502 24.600 27.348 1.00 23.85 A C ATOM 3599 CA GLN 469 33.500 28.321 1.00 20.27 A C ATOM 3599 CA GLN 469 33.712 50.158 29.899 1.00 19.50 A C ATOM 3599 CA GLN 469 33.712 50.158 29.899 1.00 19.50 A C ATOM 3600 CB GLN 470 30.410 50.136 32.961 1.00 23.85 A C ATOM 3600 CB GLN 470 30.410 50.136 32.961 1.00 23.85 A C ATOM 3600 CB GLN 470 30.410 50.136 32.961 1.00 23.85 A C ATOM 3600 CB GLN 470 30.410 50.136 32.911 1.00 23.564 A N A ATOM 3600 CB GLN 470 30.410 50.136 32.911 1.00 23.566 A				٠,		FΙ	G. 4	- 74			(Continued)
ATOM 3589 CG TYR 468 28.801 53.043 27.249 1.00 23.88 A C ATOM 3581 CDI TYR 468 25.308 53.214 27.466 1.00 23.81 A C ATOM 3582 CEI TYR 468 25.206 53.308 28.486 1.00 25.51 A C ATOM 3583 CDZ TYR 468 27.734 52.883 29.537 1.00 25.81 A C ATOM 3583 CDZ TYR 468 27.734 52.883 29.537 1.00 25.81 A C ATOM 3584 CE2 TYR 468 25.306 53.919 29.857 1.00 25.81 A C ATOM 3585 CDZ TYR 468 27.734 52.883 29.537 1.00 25.81 A C ATOM 3585 CDZ TYR 468 25.306 53.919 29.857 1.00 25.81 A C ATOM 3586 CH TYR 468 24.304 53.334 30.695 1.00 25.95 A O ATOM 3586 CH TYR 468 29.501 50.675 27.411 1.00 21.32 A C ATOM 3588 O TYR 468 28.672 50.059 28.070 1.00 22.73 A O ATOM 3589 N GLN 469 30.800 50.431 27.449 1.00 20.22 6 A N ATOM 3591 CB GLN 469 31.368 49.429 28.315 1.00 21.32 A C ATOM 3591 CB GLN 469 32.643 48.864 27.695 1.00 23.85 A C ATOM 3593 CD GLN 469 32.643 48.891 47.845 28.169 1.00 23.85 A C ATOM 3593 CD GLN 469 33.460 47.993 28.632 1.00 21.72 A C ATOM 3594 OEI GLN 469 34.891 47.845 28.169 1.00 23.85 A C ATOM 3595 CD GLN 469 33.460 47.993 28.632 1.00 21.72 A C ATOM 3595 CD GLN 469 33.480 47.993 28.632 1.00 21.72 A C ATOM 3595 CD GLN 469 35.322 46.609 27.948 1.00 23.85 A C ATOM 3596 CD GLN 469 35.322 46.609 27.948 1.00 23.85 A C ATOM 3596 CD GLN 469 35.322 46.609 27.948 1.00 23.85 A C ATOM 3596 CD GLN 469 35.322 46.609 27.948 1.00 19.50 A C ATOM 3599 CD GLN 469 35.322 46.609 27.948 1.00 19.63 A O ATOM 3599 CD GLU 470 31.602 50.03 32.001 1.00 20.27 A C ATOM 3598 CD GLU 470 31.602 50.03 32.001 1.00 20.27 A C ATOM 3600 CD GLU 470 31.602 50.03 32.001 1.00 20.27 A C ATOM 3600 CD GLU 470 32.786 49.380 32.959 1.00 19.50 A C ATOM 3600 CD LEU 470 32.768 49.380 32.959 1.00 19.97 A O ATOM 3602 CD LEU 470 32.768 49.380 32.959 1.00 19.97 A O ATOM 3601 CD ARG 471 37.91 49.814 33.610 1.00 20.17 A C ATOM 3601 CD ARG 471 37.91 49.814 33.610 1.00 20.17 A C ATOM 3601 CD ARG 471 37.91 49.814 33.610 1.00 20.17 A C ATOM 3601 CD ARG 471 37.91 49.814 33.610 1.00 20.17 A C ATOM 3611 NE ARG 471 35.927 49.386 31.261 1.00 31.73 A C ATOM 3611 NE ARG 471 35.937 49.8197 30.92 3											
ATOM 3580 CG TYR 468 27.588 53.011 28.155 1.00 24.49 A C ATOM 3581 CD1 TYR 468 26.308 53.214 27.646 1.00 23.81 A C ATOM 3582 CEI TYR 468 25.206 53.308 28.486 1.00 25.51 A C ATOM 3583 CD2 TYR 468 27.734 52.883 29.537 1.00 26.39 A C ATOM 3584 CE2 TYR 468 25.308 52.971 30.390 1.00 25.67 A C ATOM 3585 CZ TYR 468 25.308 52.971 30.390 1.00 25.67 A C ATOM 3585 CZ TYR 468 25.308 52.971 30.390 1.00 25.67 A C ATOM 3587 C TYR 468 24.304 53.334 30.695 1.00 25.95 A O ATOM 3587 C TYR 468 29.501 50.675 27.411 1.00 21.32 A C ATOM 3588 OF TYR 468 29.501 50.675 27.411 1.00 21.32 A C ATOM 3589 N GLN 469 30.800 50.431 27.449 1.00 20.26 A N ATOM 3590 CA GLN 469 31.368 49.429 28.315 1.00 19.27 A C ATOM 3591 CB GLN 469 32.643 48.864 27.695 1.00 20.12 A C ATOM 3591 CB GLN 469 33.460 47.993 28.632 1.00 20.12 A C ATOM 3593 CD GLN 469 33.460 47.993 28.632 1.00 21.72 A C ATOM 3593 CD GLN 469 33.460 47.993 28.632 1.00 21.72 A C ATOM 3595 CD GLN 469 31.712 50.158 29.589 1.00 23.85 A C ATOM 3595 CD GLN 469 31.712 50.158 29.589 1.00 23.85 A C ATOM 3595 CD GLN 469 31.712 50.158 29.589 1.00 23.84 A N A N ATOM 3596 C GLN 469 31.712 50.158 29.589 1.00 19.50 A C ATOM 3598 N LEU 470 31.277 49.113 37.794 1.00 19.63 A O ATOM 3598 N LEU 470 31.277 49.113 37.794 1.00 19.53 A C ATOM 3598 N LEU 470 31.277 49.113 37.794 1.00 19.50 A C ATOM 3600 CB LEU 470 31.477 49.113 37.794 1.00 19.50 A C ATOM 3600 CB LEU 470 31.477 49.31 33.996 1.00 19.50 A C ATOM 3600 CB LEU 470 31.477 49.31 33.996 1.00 19.977 A N A C ATOM 3600 CB LEU 470 30.40 50.42 33.84 1.00 19.977 A C ATOM 3600 CB LEU 470 30.40 50.42 33.84 1.00 20.15 A C ATOM 3600 CB LEU 470 30.40 50.42 33.84 1.00 20.15 A C ATOM 3600 CB LEU 470 30.40 50.42 33.84 1.00 20.15 A C ATOM 3600 CB LEU 470 30.40 50.42 30.40 1.00 30.58 A C A C ATOM 3600 CB LEU 470 30.40 50.40 30.40 50.40 30.00 50.50 1.00 30.58 A C A C ATOM 3600 CB LEU 470 30.20 52.60 33.84 1.00 20.15 A C A C ATOM 3600 CB LEU 470 30.40 50.40 30.40 50.50 30.00 50.40 9.977 A O A ATOM 3610 CG LEU 470 30.40 50.40 30.40 50.50 30.00 50.40 9.977 A O A C ATOM 3600 C										A	
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ATOM 3623 N SER 473 37.049 49.219 38.583 1.00 22.51 A N ATOM 3624 CA SER 473 38.377 48.809 39.022 1.00 23.17 A C											Ü
ATOM 3624 CA SER 473 38.377 48.809 39.022 1.00 23.17 A C											

				FIG. 4-75	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3649 CD: 3650 C	SER SER GLY GLY PRO PRO PRO PRO PRO GLY GLY GLY GLY GLY LEU LEU LEU LEU LEU LEU	473 473 473 474 474 474 475 475 475 475 476 476 476 477 477 477 477	39. 500 50. 976 39. 071 1. 00 23. 39 A 38. 557 48. 754 40. 536 1. 00 23. 29 A 39. 685 48. 758 41. 028 1. 00 24. 44 A 37. 457 48. 697 41. 279 1. 00 23. 29 A 37. 573 48. 627 42. 724 1. 00 23. 91 A 36. 330 49. 075 43. 459 1. 00 24. 41 A 35. 434 49. 658 42. 849 1. 00 25. 28 A 36. 257 48. 850 44. 780 1. 00 24. 58 A 35. 174 49. 389 45. 623 1. 00 25. 74 A 37. 280 48. 206 45. 609 1. 00 24. 00 A 36. 887 48. 620 47. 022 1. 00 22. 53 A 35. 419 48. 692 46. 945 1. 00 25. 59 A 37. 397 46. 692 45. 462 1. 00 24. 86 A 38. 294 46. 081 46. 044 1. 00 26. 60 A 36. 502 46. 085 44. 691 1. 00 24. 35 A 36. 502 46. 085 44. 691 1. 00 24. 35 A 36. 504 44. 646 44. 498 1. 00 23. 50 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 37. 324 44. 316 43. 227 1. 00 24. 87 A 38. 603 42. 681 41. 601 1. 00 25. 85 A 37. 324 44. 316 42. 818 1. 00 27. 45 A 38. 626 38. 851 41. 943 1. 00 27. 45 A 38. 626 38. 851 41. 943 1. 00 27. 45 A 38. 626 38. 851 41. 943 1. 00 27. 45 A	(Continued) O C O N C C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM	3651 O 3652 N 3653 CD 3654 CA	LEU PRO PRO PRO	477 477 478 478 478	97 900 40 447 40 404 4 00	
ATOM ATOM ATOM ATOM	3655 CB 3656 CG 3657 C 3658 O 3659 N	PRO PRO PRO PRO LEU	478 478 478 478 479	38. 908	C C C O N
ATOM ATOM ATOM ATOM ATOM ATOM	3661 CB 3662 CG 3663 CD1 3664 CD2	LEU LEU LEU	479 479 479 479 479	34. 376	C C C C
ATOM ATOM ATOM ATOM	3665 C 3666 O 3667 N 3668 CA 3669 CB	LEU LEU TYR TYR TYR	479 479 480 480 480	34. 077 44. 857 35. 199 1. 00 22. 18 A 33. 942 46. 073 35. 244 1. 00 22. 27 A 33. 978 44. 160 34. 073 1. 00 22. 51 A 33. 690 '44. 801 32. 790 1. 00 22. 76 A 34. 709 44. 353 31. 749 1. 00 22. 59 A	C O N C
ATOM ATOM ATOM ATOM ATOM	3672 CE1 3673 CD2	TYR TYR TYR TYR TYR	480 480 480 480 480	36. 123	C C C C

FIG. 4-77										
1.7701.4					r i d.	4 11				
ATOM	3724			487		. 464 27. 824		Α	С	
ATOM ATOM	3725 3726			487		. 229 27. 620		A	C	
ATOM	3727		1 ASN	487 487		.007 27.171		A	C	
ATOM	3728		2 ASN	487		. 948 26. 864 . 158 27. 132		A	0	
ATOM	3729		ASN.			. 220 29. 047		· A	N C	
ATOM	3730		ASN	487		. 758 30. 079		A A	0	
ATOM	3731	N	ASP	488		.514 28.929		A	N	
ATOM	3732			488	20.606 52	. 290 30. 015	1.00 23.32	Ä	Ċ	
ATOM	3733			488	20. 415 53	304 31.148		A	C	
ATOM ATOM	3734 3735	CG	ASP 1 ASP	488	20. 780 54	718 30.750		Α	C	
ATOM	3736		2 ASP	488 488		956 30.345		A	0	
ATOM	3737	C	ASP	488		601 30.862 883 30.608		A	0	
ATOM	3738	Ŏ	ASP	488		689 31.803	1.00 24.38 1.00 24.38	A	C	
ATOM	3739	N	LYS	489		902 29. 791	1.00 24.33	A A	O N	
ATOM	3740	CA	LYS	489	20.009 48.	541 30.300	1.00 25.48	A	Č	
ATOM	3741	CB	LYS	489	18.837 47.	817 29.630	1.00 25.85	Ä	č	
ATOM ATOM	3742 3743	CG	LYS	489	17. 651 47.	594 30.579	1.00 28.57	Α	C	
ATOM	3744	CD CE	LYS LYS	489 489	17. 247 48.	906 31. 251	1.00 30.67	A	C	•
ATOM	3745	NZ	LYS	489		695 32.453 944 33.278	1.00 29.68	A	C	
ATOM	3746	C	LYS	489		944 33. 278 749 30. 110	1.00 30.13 1.00 26.05	A	N	
ATOM	3747	0	LYS	489		914 29.106	1.00 26.03	A A	C 0	
ATOM	3748	N	GLY	490		894 31.084	1.00 25.12	A	N N	
ATOM	3749	CA	GLY	490	22.812 46.	094 31.019	1.00 23.91	Ä	C	
ATOM	3750	C	GLY	490		966 30.017	1.00 25.29	Ā	Č	
ATOM ATOM	3751 3752	0	GLY	490	21.855 44.		1.00 27.16	Α	0	
ATOM	3753	N CA	LEU LEU	491 491	23. 531 44.		1.00 24.58	A	N	
ATOM	3754	CB	LEU	491	23. 503 43. 24. 298 44.		1.00 24.98	A	C	
ATOM	3755	CG	LEU	491	23. 809 45.		1.00 25.21 1.00 25.03	. A	C	
ATOM	3756		LEU	491	24. 796 45.		1.00 23.03	A A	C C	
ATOM	3757	CD2	LEU	491	22. 430 45.		1.00 25.37	A	C	
ATOM	3758	C	LEU	491	24. 081 42.		1.00 25.59	Ä	Č	
ATOM	3759	0	LEU	491	23. 541 41.	579 28.250	1.00 27.45	Ä	Ŏ	
ATOM ATOM	3760 3761	N CA	ARG	492	25. 179 42.		1.00 24.68	Α	N	
ATOM	3762	CA CB	ARG ARG	492 492	25. 798 41. 1		1.00 24.07	A	C	
ATOM	3763	CG	ARG	492	26. 045 40. 5 27. 159 40. 5		1.00 24.82	A	, C	
ATOM	3764	CD	ARG	492	27. 105 40. (1.00 26.62 1.00 26.76	A	C	
ATOM	3765	NE	ARG	492	25. 884 40. 3		1.00 20.70	A A	N C	
ATOM	3766	CZ	ARG	492	25. 708 41. 4		1.00 25.45	A	C.	
ATOM	3767	NH1		492	26.684 42.2		1.00 31.57	A	N	•
ATOM		NH2		492	24.540 41.6	310 24.261	1.00 29.62	A	N	
ATOM ATOM	3769 3770	C	ARG	492	27. 117 41. 8		1.00 23.83	A	Ċ	
ATOM		0 N	ARG VAL	492 493	27.602 42.9		1.00 22.78	A	0	
ATOM		CA	VAL	493 493	27.680 40.8		1.00 24.93	A	N	
. 11 0114	3114	OIL	TAL	430	28. 966 40. 9	11 31.791	1.00 25.89	Α	C	

٠.		٠.,		FIG. 4-78	(Continued)
ATOM ATOM	3773 3774	CB V	VAL 493 VAL 493	29. 018 40. 034 33. 052 1. 00 25. 39	C C
ATOM	3775	CG2 V		27. 977 40. 482 34. 044 1. 00 25. 35 A	Ċ
ATOM	3776		VAL 493	30. 022 40. 382 30. 823 1. 00 26. 55 A	Č
ATOM	3777		VAL 493	29. 858 39. 307 30. 250 1. 00 29. 06 A	ŏ
ATOM	3778		LEU 494		Ň
ATOM	3779		LEU 494		Ċ
ATOM	3780		LEU 494	32. 657 41. 913 28. 944 1. 00 23. 74 A	Č
ATOM	3781		LEU 494	31.611 42.554 28.031 1.00 22.82 A	Č
ATOM	3782	CD1 I	LEU 494	32.017 43.989 27.697 1.00 22.34 A	С
ATOM	3783	CD2 I		31. 453 41. 706 26. 769 1. 00 19. 11 A	C C C
ATOM	3784		LEU 494	33. 315 40. 034 30. 453 1. 00 26. 29 A	
ATOM	3785		LEU 494	34. 001 39. 182 29. 885 1. 00 29. 20 A	0
ATOM	3786		GLU 495	33. 536 40. 420 31. 703 1. 00 24. 94 A	N
ATOM	3787		GLU 495	34. 623 39. 859 32. 498 1. 00 24. 93 A	C
ATOM	3788		GLU 495		C
ATOM	3789		GLU 495		C C C
ATOM ATOM	3790 3791	CD CO	GLU 495	37. 332 38. 435 32. 733 1. 00 29. 02 A	
ATOM	3792	0E1 0		37. 263 37. 724 33. 760 1. 00 29. 22 A 37. 539 37. 962 31. 596 1. 00 30. 56 A	0
ATOM	3793		LU 495		0
ATOM	3794		LU 495	34. 357 40. 210 33. 951 1. 00 25. 32 A 34. 146 41. 380 34. 285 1. 00 24. 97 A	C
ATOM	3795		ASP 496	34. 358 39. 197 34. 809 1. 00 25. 38 A	0 N
ATOM	3796		ASP 496	34. 093 39. 409 36. 224 1. 00 27. 01 A	C
ATOM	3797		ASP 496	32. 761 38. 757 36. 602 1. 00 27. 17 A	Č
ATOM	3798		ASP 496	32. 814 37. 236 36. 567 1. 00 27. 71 A	č
ATOM	3799	OD1 A		31. 755 36. 611 36. 759 1. 00 30. 85 A	ŏ
ATOM	3800	OD2 A		33. 898 36. 657 36. 360 1. 00 29. 23 A	Ö
ATOM	3801	C A	ASP 496	35. 213 38. 889 37. 127 1. 00 27. 65 A	Č
ATOM	3802		ASP 496	35. 177 39. 071 38. 345 1. 00 27. 02 A	0
ATOM	3803		ASN 497	36. 201 38. 234 36. 528 1. 00 27. 52 A	N
ATOM	3804		ISN 497	37. 329 37. 717 37. 287 1. 00 29. 40 A	C
ATOM	3805		ISN 497	38. 047 38. 863 37. 998 1. 00 28. 73 A	C
ATOM	3806		ISN 497	38. 973 39. 622 37. 080 1. 00 29. 26 A	C
ATOM	3807	OD1 A		39. 988 39. 093 36. 630 1. 00 27. 48 A	0
ATOM	3808	ND2 A		38. 628 40. 870 36. 792 1. 00 31. 42 A	N
ATOM	3809		ISN 497	36. 946 36. 652 38. 301 1. 00 30. 77 A	C
ATOM ATOM	3810 3811		ASN 497 SER 498	37. 407 36. 669 39. 444 1. 00 31. 70 A	0
ATOM	3812		SER 498 SER 498	36. 108 35. 721 37. 869 1. 00 31. 77 A 35. 666 34. 629 38. 716 1. 00 31. 32 A	N
ATOM	3813		SER 498		C
ATOM	3814		SER 498	34. 644 33. 778 37. 974 1. 00 32. 01 A 33. 520 34. 561 37. 629 1. 00 35. 01 A	C . 0
ATOM	3815		ER 498	36. 854 33. 772 39. 093 1. 00 30. 55 A	C
ATOM	3816		SER 498	37. 056 33. 456 40. 266 1. 00 31. 44 A	Ö
ATOM	3817		LA 499	37. 638 33. 398 38. 087 1. 00 29. 46 A	N
ATOM	3818		LA 499	38. 814 32. 566 38. 304 1. 00 29. 07 A	Č
ATOM	3819		LA 499	39. 626 32. 477 37. 033 1. 00 27. 47 A	č
ATOM	3820		LA 499	39. 657 33. 156 39. 421 1. 00 30. 28 A	č
ATOM	3821	0 · A	LA 499	39. 885 32. 515 40. 447 1. 00 30. 98 A	0

	(Continued)					
ATOM ATOM ATOM	3822 3823 3824	N CA CB	LEU LEU LEU	500 500 500	F I G. 4 - 7 9 40.098 34.393 39.223 1.00 30.98 A 40.919 35.073 40.208 1.00 31.89 A 41.218 36.502 39.755 1.00 31.32 A	N C C
ATOM ATOM ATOM	3825 3826 3827		LEU LEU. LEU	500 500 500	42. 106 37. 312 40. 703 1. 00 31. 18 A 43. 459 36. 635 40. 871 1. 00 29. 43 A 42. 269 38. 711 40. 155 1. 00 31. 85 A	C C C
ATOM ATOM ATOM	3828 3829	C 0	LEU LEU	500 500	40. 251 35. 096 41. 574 1. 00 33. 26 A 40. 878 34. 772 42. 578 1. 00 33. 38 A	C 0
ATOM ATOM	3830 3831 3832	N CA CB	ASP ASP ASP	501 501 501	38. 984 35. 484 41. 624 1. 00 35. 48 A 38. 294 35. 522 42. 905 1. 00 38. 46 A 36. 815 35. 859 42. 720 1. 00 40. 04 A	N C .C
ATOM ATOM ATOM	3833 3834 3835	CG OD1 OD2	ASP ASP ASP	501 501 501	36. 068 35. 942 44. 043 1. 00 42. 67 A 36. 349 36. 870 44. 831 1. 00 44. 51 A 35. 202 35. 076 44. 300 1. 00 44. 58 A	C 0 0
ATOM ATOM ATOM	3836 3837 3838	C O N	ASP ASP LYS	501 501 502	38. 432 34. 149 43. 557 1. 00 39. 76 A 38. 622 34. 039 44. 765 1. 00 39. 03 A 38. 352 33. 103 42. 740 1. 00 41. 28 A	C O N
ATOM ATOM ATOM	3839 3840 3841	CA CB CG	LYS LYS LYS	502 502 502	38. 470 31. 741 43. 237 1. 00 42. 62 A 38. 206 30. 746 42. 100 1. 00 44. 22 A	C C C
ATOM ATOM ATOM	3842 3843 3844	CD CE NZ	LYS LYS LYS	502 502 502	39. 071 28. 557 43. 050 1. 00 47. 22 A 38. 700 27. 147 43. 516 1. 00 47. 98 A	C C
ATOM ATOM	3845 3846	C 0	LYS LYS	502 502	39. 866 31. 534 43. 828 1. 00 43. 11 A 40. 001 31. 079 44. 963 1. 00 43. 40 A	N C O
ATOM ATOM ATOM	3847 3848 3849	N CA CB	MET MET MET	503 503 503	40. 900 31. 881 43. 064 1. 00 42. 72 A 42. 280 31. 735 43. 528 1. 00 43. 17 A 43. 256 32. 193 42. 444 1. 00 45. 35 A	N C C
ATOM ATOM ATOM	3850 3851 3852	CG SD CE	MET MET MET	503 503 503	43. 267 31. 332 41. 200 1. 00 48. 35 A 44. 396 32. 004 39. 952 1. 00 54. 36 A 45. 957 31. 226 40. 438 1. 00 52. 89 A	C S C
ATOM ATOM ATOM	3853 3854 3855	C O N	MET MET LEU	503 503 504	42.551 32.530 44.807 1.00 41.81 A 43.059 31.990 45.790 1.00 40.44 A 42.215 33.815 44.779 1.00 41.12 A	C O N
ATOM ATOM ATOM	3856 3857 3858	CA CB CG	LEU LEU LEU	504 504 504	42. 412 34. 700 45. 919 1. 00 42. 37 A 41. 914 36. 103 45. 566 1. 00 41. 90 A	C C
ATOM ATOM ATOM	3859 3860 3861	CD1 CD2	LEU LEU	504 504	44. 111 36. 668 44. 472 1. 00 41. 70 A 42. 277 38. 376 44. 635 1. 00 40. 64 A	C C C
ATOM ATOM	3862 3863	C O N	LEU LEU GLN	504 504 505	41. 727 34. 211 47. 199 1. 00 43. 78 A 42. 056 34. 664 48. 298 1. 00 43. 47 A 40. 774 33. 292 47. 054 1. 00 44. 74 A	C O N
ATOM ATOM ATOM	3864 3865 3866	CA CB CG	GLN GLN GLN	505 505 505	40. 053 32. 737 48. 198 1. 00 45. 12 A 38. 911 31. 834 47. 721 1. 00 47. 10 A 37. 767 32. 574 47. 059 1. 00 50. 85 A	C C C
ATOM ATOM ATOM	3867 3868 3869	CD OE1 NE2	GLN GLN GLN	505 505 505	37. 091 33. 544 48. 005 1. 00 52. 28 A 36. 320 33. 143 48. 878 1. 00 53. 91 A 37. 390 34. 829 47. 848 1. 00 53. 20 A	C O N
ATOM	3870	C	GLN	505	40. 981 31. 920 49. 090 1. 00 44. 28 A	C

	(3) 2 ••			FΙ	G. 4		(Continued)		
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3871	GLN ASN ASN ASN ASN ASN ASN VAL VAL VAL VAL VAL CLN GLN GLN GLN GLN	505 506 506 506 506 506 506 507 507 507 507 507 507 507 508 508 508	F I 40. 806 41. 970 42. 907 43. 301 43. 962 44. 478 43. 945 44. 156 45. 191 44. 060 45. 186 45. 801 46. 989 46. 234 44. 726 43. 617 45. 586 45. 272 46. 146 46. 034 46. 955	31. 863 31. 288 30. 452 29. 254 28. 157 27. 187 28. 293 31. 211 30. 605 32. 538 33. 367 34. 155 34. 974 33. 204 34. 369 34. 887 34. 634 35. 578 35. 307 33. 894 33. 667		1. 00 44. 07 1. 00 43. 04 1. 00 43. 10 1. 00 47. 04 1. 00 50. 97 1. 00 52. 33 1. 00 41. 53 1. 00 41. 53 1. 00 39. 25 1. 00 35. 74 1. 00 35. 80 1. 00 34. 07 1. 00 34. 07 1. 00 33. 19 1. 00 31. 62 1. 00 31. 59 1. 00 30. 69	A A A A A A A A A A A A A A A A A A A	(Continued) 0 N C C C O N C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3892 OE1	GLN GLN GLN MET MET MET MET MET MET MET PRO	508 508 508 508 509 509 509 509 509 510 510 510 511 511 511 511 512 512	46. 994 47. 696 45. 521 46. 480 44. 652 44. 775 43. 744 44. 004 45. 605 45. 130 44. 602 43. 875 45. 279	34. 471 32. 568 36. 996 37. 648 37. 463 38. 797 38. 993 38. 143 38. 540 39. 727 39. 890 39. 724 41. 032 41. 361 42. 985 42. 847 42. 886 42. 632 43. 826 44. 592 44. 441 44. 897 46. 043 46. 433 46. 838 48. 242 48. 555	57. 083 56. 130 52. 689 53. 097 51. 801 51. 236 50. 124 48. 896 48. 171 46. 922 52. 280 53. 255 52. 085	1.00 30.03 1.00 31.83 1.00 28.80 1.00 30.18 1.00 29.60 1.00 28.77 1.00 28.64 1.00 30.06 1.00 31.71 1.00 34.08 1.00 27.67 1.00 28.41 1.00 26.51 1.00 25.01 1.00 24.17 1.00 24.17 1.00 23.17 1.00 23.17 1.00 24.30 1.00 23.52 1.00 23.52 1.00 21.47 1.00 25.81 1.00 27.50 1.00 27.50 1.00 27.50 1.00 24.17 1.00 25.44 1.00 23.04	A A A A A A A A A A A A A A A A A A A	O N C O N C C C S C C O N

					E 1 (~ 4	0 1			(Continued)
					r I (G. 4	- 8 1			
ATOM ATOM ATOM	3920 3921 3922	CG CD CE	LYS LYS LYS	512 512 512	42. 252 42. 368 42. 639	50. 043 50. 249 51. 688	50. 621 49. 125 48. 792	1.00 21.12 1.00 21.07 1.00 19.46	Α	C C C
ATOM ATOM	3923 3924	NZ C	LYS LYS	512 512	42. 779 41. 095	51.870 49.109	47. 343 53. 105	1.00 15.68 1.00 24.25	A A	N C
ATOM ATOM	3925 3926	0 N	LYS LYS	512 513	39. 905 41. 546	48. 958 50. 017	52.846	1.00 23.45 1.00 24.50	Α	0
ATOM	3927	CA	LYS	513	40.661	50.941	53. 960 54. 647	1.00 25.28	A A	N C
ATOM ATOM	3928 3929	CB CG	LYS LYS	513 513	41.040 40.202	51. 041 52. 025	56. 124 56. 914	1.00 26.65 1.00 27.55	A A	C C C C
ATOM ATOM	3930 3931	CD CE	LYS LYS	513 513	38. 754	51.577 52.476	56.954	1.00 33.11	Α	Č
ATOM	3932	NZ	LYS	513	36. 503	51.943	57. 844 57. 960	1.00 35.12 1.00 38.12	A A	C N
ATOM ATOM	3933 3934	C 0	LYS LYS	513 513		52. 312 52. 829	53. 999 53. 877	1.00 26.42 1.00 28.66	A A	0 C ·
ATOM ATOM	3935 3936	N CA	LEU LEU	514 514	39.688	52. 891 54. 213	53.575	1.00 25.40	Α	N
ATOM	3937	CB	LEU	514	39. 147	54.119	52.958 51.536	1.00 22.53 1.00 20.88	A A	C C
ATOM ATOM	3938 3939	CG CD1	LEU LEU	514 514		55. 443 56. 242	50. 825 50. 662	1.00 21.52 1.00 20.94	A A	C C
ATOM ATOM	3940 3941		LEU LEU	514 514	38. 244	55. 153 55. 151	49. 476 53. 788	1.00 22.59	Α	C
ATOM	3942	0	LEU	514	37. 591	54. 981	53.844	1.00 22.73 1.00 20.65	A A	C 0
ATOM ATOM	3943 3944	N CA	ASP ASP	515 515		56. 132 57. 076	54. 437 55. 268	1.00 23.05 1.00 25.43	A A	N C
ATOM ATOM	3945 3946	CB CG	ASP ASP	515 515	38. 581	56. 535 57. 142	56. 693 57. 458	1.00 27.35	Α	C
ATOM	3947	0D1	ASP	515	37. 278	56.851	58.668	1.00 30.82 1.00 32.73	A A	C 0
ATOM ATOM	3948 3949	OD2 C	ASP ASP	515 515		57. 905 58. 462	56. 851 55. 287	1.00 32.89 1.00 26.80	A A	C 0
ATOM ATOM	3950 3951	0 N	ASP PHE	515	40.054	58. 835	54.357	1.00 27.23	Α	0
ATOM	3952	CA	PHE	516 516	39.688	59. 230 60. 566	56. 345 56. 431	1.00 27.53 1.00 28.71	A A	N C
ATOM ATOM	3953 3954	CB CG	PHE PHE	516 516		61.590 61.658	55. 729 56. 291	1.00 28.60 1.00 28.84	A A	C C
ATOM ATOM	3955 3956	CD1 CD2	PHE	516	37.160	62.115	57. 583	1.00 29.59	Α	С
ATOM	3957	CE1	PHE	516 516	35.875	61. 242 62. 157	55. 532 58. 116	1.00 30.94 1.00 28.99	A A	C C
ATOM ATOM	3958 3959	CE2 CZ	PHE PHE	516 516		61. 279 61. 737	56. 058 57. 352	1.00 29.88 1.00 29.33	A A	C C
ATOM ATOM	3960 3961	C 0	PHE PHE	516	39. 943	61.024	57.861	1.00 28.58	Α	C
ATOM	3962	N	ILE	516 517	40.773	60. 450 62. 053	58. 811 57. 990	1.00 29.42 1.00 26.80	A· A	O N
ATOM ATOM	3963 3964	CA CB	ILE ILE	517 517		62. 651 62. 410	59. 272 59. 686	1.00 28.68 1.00 27.66	A A	C
ATOM	3965	CG2	ILE	517	42.799	60. 937	59.989	1.00 23.78	Α	C
ATOM ATOM	3966 3967	CG1 CD1		517 517		62. 861 64. 361	58. 581 58. 431	1.00 29.30 1.00 31.79	A A	C C
ATOM	3968	C	ILE	517		64.132	59.041	1.00 30.84	A	č

					(Continued)
				FIG. 4-82	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	3969 3970 3971 3972 3973 3974 3975 3976 3977 3978 3979 3980 3981	O ILE N ILE CA ILE CB ILE CG1 ILE CD1 ILE O ILE N LEU CA LEU CG LEU CD1 LEU	519 519 519 519 519	40. 813 64. 577 57. 898 1. 00 31. 70 A 40. 616 64. 899 60. 102 1. 00 32. 28 A 40. 323 66. 313 59. 924 1. 00 33. 51 A 38. 977 66. 683 60. 595 1. 00 33. 41 A 38. 603 68. 125 60. 283 1. 00 33. 29 A 37. 871 65. 765 60. 072 1. 00 33. 38 A 36. 535 65. 972 60. 749 1. 00 33. 46 A 41. 415 67. 222 60. 455 1. 00 35. 00 A 41. 883 67. 069 61. 580 1. 00 35. 82 A 41. 824 68. 169 59. 622 1. 00 36. 74 A 42. 850 69. 126 59. 997 1. 00 39. 19 A 44. 169 68. 828 59. 276 1. 00 38. 52 A 44. 746 67. 413 59. 364 1. 00 39. 31 A	N C C C C C C O N C C C C
ATOM ATOM	3983 3984	CD2 LEU C LEU	519 519	45.068 67.059 60.806 1.00 39.59 A 42.351 70.501 59.591 1.00 40.26 A	C
ATOM ATOM ATOM ATOM	3985 3986 3987 3988	O LEU N ASN CA ASN CB ASN	519 520 520 520	42. 102 70. 754 58. 414 1. 00 40. 93 A 42. 198 71. 382 60. 574 1. 00 41. 70 A 41. 736 72. 735 60. 321 1. 00 42. 46 A 42. 760 73. 474 59. 467 1. 00 44. 27 A	O N C
ATOM	3989	CG ASN	520	44. 078 73. 635 60. 177 1. 00 46. 04 A	C .
ATOM ATOM	3990 3991	OD1 ASN ND2 ASN	520 520	44. 540 72. 723 60. 859 1. 00 47. 21 A 44. 697 74. 796 60. 020 1. 00 50. 39 A	
ATOM ATOM	3992 3993	C ASN O ASN	520 520	40. 384 72. 728 59. 638 1. 00 42. 18 A 40. 183 73. 388 58. 620 1. 00 42. 15 A	
ATOM ATOM	3994 3995	N GLU	521	39.461 71.963 60.210 1.00 41.73 A	N
ATOM	3996	CA GLU CB GLU	521 521	38. 105 71. 861 59. 691 1. 00 42. 64 A 37. 445 73. 245 59. 660 1. 00 44. 72 A	
ATOM ATOM	3997 3998	CG GLU	521 521	37. 967 74. 204 60. 715 1. 00 48. 09 A 38. 057 73. 564 62. 081 1. 00 50. 91 A	C C
ATOM	3999	OE1 GLU	521	36.994 73.245 62.661 1.00 52.95 A	0
ATOM ATOM	4000 4001	OE2 GLU C GLU	521 521	39. 194 73. 374 62. 568 1. 00 51. 94 A 38. 041 71. 248 58. 296 1. 00 40. 90 A	0 C
ATOM ATOM	4002 4003	O GLU N THR	521 522	36. 967 71. 171 57. 701 1. 00 40. 88 A	0
ATOM	4004	CA THR	522	39. 182 70. 814 57. 772 1. 00 39. 01 A 39. 206 70. 221 56. 442 1. 00 36. 94 A	N C
ATOM ATOM	4005 4006	CB THR OG1 THR	522 522	40. 339 70. 816 55. 584 1. 00 38. 55 A 40. 127 72. 223 55. 431 1. 00 40. 51 A	C 0
ATOM ATOM	4007 4008	CG2 THR C THR	522 522	40. 364 70. 171 54. 202 1. 00 39. 39 A	С
ATOM	4009	0 THR	522	40.086 68.152 57.305 1.00 33.48 A	C 0
ATOM ATOM	4010 4011	N LYS CA LYS	523 523	38. 653 68. 045 55. 573 1. 00 33. 07 A 38. 685 66. 597 55. 479 1. 00 30. 63 A	N C
ATOM	4012	CB LYS	523	37. 357 66. 105 54. 901 1. 00 31. 78 A	С
ATOM ATOM	4013 4014	CG LYS	523 523	36. 882 64. 770 55. 440 1. 00 34. 92 A 35. 473 64. 458 54. 956 1. 00 37. 12 A	C C
ATOM	4015 4016	CE LYS NZ LYS	523 523	34. 473 65. 488 55. 455 1. 00 40. 20 A	C
ATOM ATOM	4017	C LYS	523	33. 111 65. 296 54. 873 1. 00 43. 74 A 39. 845 66. 191 54. 576 1. 00 28. 84 A	N C

										•		(Con	tinuad)
					F]	G.	4	- 83				(Con	tinued)
		_											
ATOM	4018	0	LYS	523	39. 962		661	53. 448		29.90	A	0	
ATOM	4019	N	PHE	524	40.711		329	55.086		26.11	A	N	
ATOM	4020	CA	PHE	524	41.857		858	54. 334		23. 17	A	C	
ATOM	4021	CB	PHE	524	43. 139		407	54. 953		22.95	A	C	
ATOM	4022	CG	PHE	524	43. 394		854	54.636		21. 35	A	C	
ATOM	4023		PHE	524	43. 773		242	53. 346		21.14	A	C	
ATOM	4024		PHE	524	43. 265		830	55. 620		18.86	A	C	
ATOM	4025		PHE	524	44.026		587	53.040		19. 22	Ą	C	
ATOM	4026		PHE	524	43. 512			55. 329		19.37	A	C	
ATOM	4027	CZ	PHE	524	43. 895		552	54. 034		19.34	A	C	
ATOM	4028	C	PHE	524	41.872		337	54. 328		23. 15	A	C	
ATOM	4029	0	PHE	524	42.084		703	55. 356		22.01	Α.	0	
ATOM	4030	N	TRP	525	41.640		758	53. 156		24.00	A	N	
ATOM	4031	CA	TRP	525	41.593		309	53.000		23.65	A	C	
ATOM	4032	CB	TRP	525	40.875		958	51.696		23. 74	A	C	
ATOM	4033	CC	TRP	525 525	39.476			51.647		24. 69	A	C	
ATOM	4034	CD2	TRP TRP	525	38. 291			51.893		25. 25	A	C .	
ATOM ATOM	4035 4036		TRP	525	37. 195			51.800		26.02	A	C	
ATOM	4030		TRP	525 525	38. 049			52. 186		25. 53	A	C	
ATOM	4038		TRP	525 525	39.065			51.418		25. 58	A	C	
ATOM	4039		TRP	525 525	37. 693 35. 874			51. 508 51. 990		25. 32	A	N	
ATOM	4040		TRP	525 525	36. 735			51. 990		25. 72 24. 54	A	C	
ATOM	4041	CH2		525	35. 666			52. 276		24. 86	A	C	
ATOM	4042	C	TRP	525	42. 927			53. 042		23. 39	A	C	
ATOM	4043	Õ	TRP	525	43. 994			52. 803		24. 19	Α.	0	
ATOM	4044	N	TYR	526	42. 840			53. 347		22. 63	A A	N	
ATOM	4045	CA	TYR	526	44. 002			53. 410		22. 38	A	C	
ATOM	4046	CB	TYR	526	44. 715			54. 763		22. 15	A	C	
ATOM	4047	CG	TYR	526	43.946			55. 929		24. 08	A	C	
ATOM	4048	CD1	TYR	526	43.968			56. 178		23. 01	A	Č	
ATOM	4049	CE 1		526	43. 215			57. 204		25. 01	A	C	
ATOM	4050	CD2		526	43.150			56. 747		24. 62	A	Č	
ATOM	4051		TYR	526	42. 395			57. 772°		24. 74	A	Č	
ATOM	4052	CZ	TYR	526	42.426			57. 997		25. 67	Ä	Č	
ATOM	4053	OH	TYR	526	41.650			59.003		25. 43	A	ŏ	
ATOM	4054	C	TYR	526	43. 478			53. 251		22.00	· A	Č	
ATOM	4055	0	TYR	526	42. 294			53. 482		21.71	A	ŏ	
ATOM	4056	N	GLN	527	44.353			52. 843		19.68	A	Ň	
ATOM	4057	CA	GLN	527	43.964			52. 707		20.14	Ä	Ċ	
ATOM	4058	CB	GLN	527	43.842			51. 238		19.56	A	č	
ATOM	4059	CG	GLN	527	45.123			50.465		23.06	A	Č	
ATOM	4060	CD	GLN	527	44. 986			49.065		23. 49	Â	č	
ATOM	4061	0E1		527	44.034			48. 359		25. 79	Ä	ŏ	
ATOM	4062		GLN	527	45. 937			48.648		22.35	A	Ň	
ATOM	4063	C	GLN	527	45.038			53.389		20.67	A	Č	
ATOM	4064	0	GLN	527	46.172			53.563	1.00		Ä	Ö	
ATOM	4065	N	MET	528	44.674			53.792	1.00		A	N	
ATOM	4066	CA	MET	528	45.610		771	54.460	1.00	22.32	Α	.C	

	••.					,				(Continued)
	•				FΙ	G. 4	- 84			
ATOM	4067	СВ	MET	52 8	45.372	51.753	55.967	1.00 23.57	Α	С
ATOM	4068	CG	MET	528	45.830	52.971	56.727	1.00 23.53	Ä	Ċ
ATOM	4069	SD	MET	528	45.605	52. 683	58. 492	1.00 23.56	Ä	Š
ATOM	4070	CE	MET	528	46. 400	54. 107	59. 158	1.00 21.91	Ä	č
ATOM	4071	C	MET	528	45. 482	50. 347	53. 974	1.00 23.25	Ä	Č
ATOM	4072	Õ	MET	528	44. 383	49. 790	53. 935	1.00 24.82	A	Ö
ATOM	4073	N	ILE	529	46.605	49. 751	53.600	1.00 22.51	Ä	N
ATOM	4074	CA	ILE	529	46. 587	48. 363	53. 183	1.00 21.97	Ä	Ċ ·
ATOM	4075	CB	ILE	529	47.644	48.078	52. 116	1.00 19.54	Ä	č
ATOM	4076		ILE	529	47. 557	46.635	51.681	1.00 13.04	Ä	Č
ATOM	4077		ILE	529	47. 454	49.029	50. 927	1.00 21.01	A	Č
ATOM	4078		ILE	529	46.045	49. 038	50. 335	1.00 19.28	A	Č.
ATOM	4079	CDI	ILE	529	46. 937	47. 620	54. 465	1.00 24.02	A	č
	4019	0	ILE	529	48. 114	47. 505	54. 820	1.00 25.51	A	Ŏ
ATOM	4080	N	LEU	52 <i>5</i> 530	45. 911	47. 153	55. 175	1.00 24.47	A	N
ATOM		CA	LEU	530 530	46.114	46. 443	56. 438	1.00 24.41	A	C
ATOM	4082			530 530	44.915	46. 640	57. 370	1.00 24.70	A	Č
ATOM	4083	CB CG	LEU	530 530	44. 451	48. 052	57. 726	1.00 24.00		C
ATOM	4084		LEU	530 530	43. 365	47. 928	58. 763	1.00 24.92	A	C
ATOM	4085		LEU					1.00 25.70	A	
ATOM	4086		LEU	530	45.589	48.896	58. 272		A	C .
ATOM	4087	C	LEU.	530 530	46.337	44. 953	56.241	1.00 24.39 1.00 24.58	A	
ATOM	4088	0	LEU	530	45.686	44.319	55.411		A	0 N
ATOM	4089	N	PRO	531	47. 272	44.374	57.003	1.00 24.58	A	N C
ATOM	4090	CD	PRO	531	48.174	45.045	57. 950	1.00 24.42	A	C
ATOM	4091	CA	PRO	531	47.578	42.943	56. 913	1.00 26.79	A	C
ATOM	.4092	CB	PRO	531	48. 763	42. 784	57. 862	1.00 26.36	A	C
ATOM	4093	CG	PRO	531	48. 580	43. 913	58. 838	1.00 26.79	A	C
ATOM	4094	C	PRO	531	46. 388	42.078	57. 312	1.00 28.05	A	C
ATOM	4095	0	PRO	531	45.443	42.562	57. 931	1.00 31.01	A	0
ATOM	4096	N	PRO	532	46.417	40. 782	56. 964	1.00 28.42	A	N
ATOM	4097	CD	PRO	532	47. 484	40.062	56. 253	1.00 28.00	A	C
ATOM	4098	CA	PRO	532	45. 316	39.874	57. 306	1.00 28.68	A	C
ATOM	4099	CB	PRO	532	45. 783	38. 534	56. 745	1.00 28.68	A	C .
ATOM	4100	CG	PRO	532	46. 726	38. 912	55.659	1.00 28.50	A	C
ATOM	4101	C	PRO	532	45. 113	39. 799	58. 814	1.00 29.80	A	C
ATOM	4102	0	PRO	532	46.051	40.006	59. 579	1.00 31.52	A	0
ATOM	4103	N	HIS	533	43.894	39.501	59. 242	1.00 31.29	A	N
ATOM	4104	CA	HIS	533	43.605	39. 382	60.670	1.00 31.80	A	C
ATOM	4105	CB	HIS	533	44. 278	38. 127	61. 225	1.00 29.82	A	C
ATOM	4106	CG	HIS	533	44. 170	36. 936	60. 324	1.00 29.23	A	. Č
ATOM	4107		HIS	533	45.114	36. 247	59.641	1.00 28.40	A	C
ATOM	4108		HIS	533	42.966	36. 335	60.024	1.00 28.40	A	N ·
ATOM	4109		HIS	533	43. 174	35. 326	59. 197	1.00 28.67	A	C
ATOM	4110		HIS	533	44. 469	35. 251	58.949	1.00 28.85	A	N
ATOM	4111	C	HIS	533	44.101	40.601	61.445	1.00 33.77	A	С
ATOM	4112	0	HIS	533	44.469	40.489	62.617	1.00 33.99	Α	0
ATOM	4113	N	PHE	534	44. 121	41.758	60.787	1.00 35.52	Α	N
ATOM	4114	CA	PHE	534	44.578	42.987	61.427	1.00 37.29	Α	С
ATOM	4115	CB	PHE	534	44. 249	44. 203	60. 555	1.00 36.11	A	C

					FIG. 4-85		(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4116 4117 4118 4120 4121 4122 4123 4124 4125 4126 4131 4133 4133 4134 4135 4136 4140 4141 4142 4143 4144 4144 4145 4145 4145 4146 4147 4147 4148 4149 4151 4151 4151 4151 4151 4151 4151	CD2 CE1 CE2 CZ C O N CA CB CG OD1 CA CB CC	PHE PHE PHE ASP ASP ASP ASP ASP LYS LYS LYS LYS LYS SER SER LYS	534 534 534 534 534 535 535 535 535 535	44. 510 45. 523 61. 235 1. 00 35. 46 45. 811 45. 956 61. 475 1. 00 35. 65 43. 455 46. 320 61. 654 1. 00 33. 35 46. 056 47. 167 62. 124 1. 00 36. 55 43. 688 47. 530 62. 304 1. 00 35. 26 44. 990 47. 957 62. 541 1. 00 35. 35 43. 920 43. 158 62. 790 1. 00 38. 07 42. 705 43. 046 62. 911 1. 00 38. 83 44. 725 43. 435 63. 810 1. 00 39. 27 44. 206 43. 621 65. 160 1. 00 40. 72 44. 751 42. 541 66. 089 1. 00 40. 72 44. 751 42. 571 67. 460 1. 00 46. 19 43. 704 43. 668 67. 912 1. 00 46. 58 43. 999 41. 499 68. 092 1. 00 40. 91 45. 799 45. 270 65. 837 1. 00 40. 57 43. 635 45. 822 66.	A A A A A A A A A A A A A A A A A A A	CCCCCCONCCCOOCONCCCCCNCONCCOCONCCCCCNC
ATOM ATOM ATOM ATOM	4155 4156 4157 4158	O N CA CB	LYS LYS LYS LYS	538 539 539 539	48. 526 47. 229 67. 283 1. 00 38. 17 50. 601 46. 485 67. 725 1. 00 36. 92 51. 263 47. 629 67. 116 1. 00 36. 43 52. 293 48. 225 68. 079 1. 00 37. 32	A A A	O N C C
ATOM ATOM ATOM ATOM ATOM	4159 4160 4161 4162 4163	CG CD CE NZ C	LYS LYS LYS LYS LYS	539 539 539 539 539	51. 693 48. 838 69. 341 1. 00 37. 42 50. 925 50. 117 69. 028 1. 00 40. 01 50. 209 50. 674 70. 258 1. 00 41. 64 51. 121 51. 014 71. 389 1. 00 43. 98 51. 943 47. 110 65. 849 1. 00 35. 38	A A A A	C C C N C
ATOM	4164	0	LYS	539	52. 699 46. 137 65. 893 1. 00 35. 49	A	0

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(Continued) FIG. 4-86 51.658 47.747 64.719 1.00 33.00 N **ATOM** 4165 N TYR 540 52.229 47.316 63.452 1.00 30.12 A C CA 540 4166 TYR **ATOM** 47.135 62.397 1.00 28.99 C 51.131 Α 4167 CB TYR 540 **ATOM** C 50.204 62.630 1.00 29.13 540 45.968 Α 4168 CG TYR **ATOM** 49.109 1.00 28.32 63.488 46.078 **ATOM** 4169 CD1 TYR 540 C 48.254 63.699 1.00 27.13 540 45.000 Α **ATOM** 4170 CE1 TYR 50.421 44.748 61.990 1.00 27.62 Α **ATOM** 4171 CD2 TYR 540 C CE2 TYR 540 49.576 43.669 62.196 1.00 26.32 A 4172 **ATOM** 1.00 27.64 C 63.051 CZ 48.495 43.800 Α 4173 TYR 540 **ATOM** 42.724 63.260 1.00 29.67 0 ОН 47.661 A ATOM 4174 TYR 540 1.00 29.33 C **ATOM** 4175 C TYR 540 53.242 48. 287 62.890 Α 53.130 540 49.492 63.091 1.00 31.23 0 **ATOM** 4176 0 TYR Α 54.270 62.199 1.00 27.71 4177 N PR₀ 541 47.772 Α N **ATOM** 54.717 46.383 62.020 1.00 25.95 A C 4178 PRO 541 **ATOM** CD 61.634 C 4179 CA PRO 55.238 48.708 1.00 27.56 A **ATOM** 541 C 4180 56.361 47.794 61.148 1.00 26.81 CB **ATOM** PR₀ 541 Α Ċ 60.867 1.00 25.92 ATOM CG PR0 541 55.662 46.512 Α 4181 \mathbf{c} 54.463 49.358 60.500 1.00 27.83 C 4182 PR₀ Α ATOM 541 4183 53.579 48.727 59.912 1.00 28.03 0 **ATOM** 0 PR₀ 541 A 4184 N 542 54.763 50.613 60.200 1.00 27.70 **ATOM** LEU Α N 542 54.032 51.307 59.154 1.00 26.55 \mathbf{c} **ATOM** 4185 CA LEU A 53.220 52.440 C 542 59.791 1.00 26.11 ATOM 4186 CB LEU Α Ċ ATOM 4187 CG LEU 542 52.252 53.292 58.959 1.00 28.68 A 59.898 C 4188 CD1 LEU 542 51.422 54.170 1.00 29.38 **ATOM** Α Ċ 4189 1.00 29.52 CD2 LEU 542 53.017 54.165 57.979 ATOM Α 54. 924 1.00 26.16 **ATOM** 4190 C LEU 542 51.855 58.042 A C 4191 0 LEU 542 55.943 52.492 58.303 1.00 28.00 0 ATOM A 54.536 51.589 56.801 1.00 23.70 4192 N LEU 543 **ATOM** Α N 55.651 **ATOM** 4193 CA LEU 543 55.263 52.097 1.00 24.11 A C 1.00 24.05 C 4194 CB LEU 543 55.595 50.978 54.660 ATOM Α 1.00 22.45 CG LEU 543 56.080 51.474 53. 289 C ATOM 4195 A 57.209 4196 53.475 C **ATOM** CD1 LEU 543 52.487 1.00 24.00 A CD2 LEU 56.537 50.303 1.00 20.16 C ATOM 4197 543 52.441 A C C 54.966 **ATOM** 4198 LEU 543 54.378 53.131 1.00 24.37 Α **ATOM** 4199 0 LEU 543 53.283 52.819 54.511 1.00 25.72 0 Α 4200 N 544 54.857 54.362 54.896 1.00 24.80 **ATOM** LEU Α N 4201 CA LEU 544 54.098 55.436 54.278 1.00 23.74 A C **ATOM** 1.00 23.92 C **ATOM** 4202 CB LEU 544 54.424 56.757 54.979 A 54.581 C **ATOM** 4203 CG LEU 544 53.640 58.003 1.00 22.62 A 1.00 24.91 Č ATOM 4204 CD1 LEU 544 52.157 57.743 54.729 A 54.069 1.00 24.25 C **ATOM** 4205 CD2 LEU 544 59.166 55.460 Α 1.00 23.24 **ATOM** 4206 C LEU 544 54.403 55.543 52.785 C Α **ATOM** 4207 0 LEU 544 55.451 56.053 52.400 1.00 23.44 A 0 1.00 21.43 **ATOM** 4208 N **ASP** 545 53.477 55.049 51.962 N A 1.00 20.10 **ATOM** 4209 CA ASP 545 53.595 55.075 50.508 Α C ASP 545 52.570 49.902 1.00 20.20 C ATOM 4210 CB 54.132 Α 1.00 20.73 C **ASP** 48.444 CG 545 53.848 ATOM 4211 52.826 Α 1.00 22.69 47.699 OD1 ASP 545 54.790 Α 0 ATOM 4212 53.175 OD2 ASP 545 1.00 19.91 Α 0 ATOM 4213 52.660 52.675 48.044

					FΙ	G. 4	- 88			(Continue
ATOM	4263	N	GLN	553	46.632	61.885	43. 926	1.00 17.07	A	N
ATOM	4264	CA	GLN	553	45.628	61.179	44.699	1.00 16.87	Ā	C
ATOM	4265	CB	GLN	553	44.301	61.090	43.937	1.00 16.43	Ä	Č
ATOM	4266	CG	GLN	553	43. 249	60. 292	44.695	1.00 19.53	Ä	Č
ATOM	4267	CD	GLN	553	41.844	60. 468	44. 163	1.00 18.87	Ä	Č
ATOM	4268	0E1		553	41.520	60. 019	43.066	1.00 20.67	Ä	Ö
ATOM	4269			553	40. 999	61.126	44.944	1.00 18.67	A	Ň
ATOM	4270	C	GLN	553	46.123	59. 781	44.996	1.00 18.09	Ä	Ċ
ATOM	4271	Ō	GLN	553	46.088	58. 915	44.129	1.00 18.25	Ä	Ö
ATOM	4272	N	LYS	554	46.589	59. 562	46. 221	1.00 19.53	Ā	N
ATOM	4273	CA	LYS	554	47.075	58. 248	46.620	1.00 20.69	Ä	Ċ
ATOM	4274	CB	LYS		48.319	58. 387	47.490	1.00 22.65	Ā	Ċ
ATOM	4275	CG	LYS	554	49.538	58.887	46.733	1.00 24.15	A	Č
ATOM	4276	CD	LYS	554	50.064	57.840	45.765	1.00 25.21	Ā	Ċ
ATOM	4277	CE	LYS	554	50.777	56. 711	46.503	1.00 24.75	A	Ċ
ATOM	4278	NZ	LYS	554	51.472	55. 796	45.560	1.00 23.89	Α	N
ATOM	4279	C	LYS	554	45.996	57.472	47.374	1.00 21.48	Α	С
ATOM	4280	0	LYS	554	46.108	56. 258	47. 549	1.00 22.39	A	0
ATOM	4281	N	ALA	555	44.952	58. 176	47.807	1.00 20.77	A	N
ATOM	4282	CA	ALA	555	43.849	57. 555	48. 538	1.00 20.46	Α	C
ATOM	4283	CB	ALA	555		58. 376	49.768	1.00 18.05	Α	C
ATOM	4284	C	ALA	555	42.611	57.436	47.643	1.00 21.32	Α	C
ATOM	4285	0	ALA	555	41.996	58. 442	47. 285	1.00 21.75	Α	0
ATOM	4286	N	ASP	556	42. 249	56. 208	47. 283	1.00 21.00	A	N
ATOM	4287	CA	ASP	556	41.096	55. 981	46.419	1.00 20.04	A	Ċ
ATOM	4288	CB	ASP	556	41.500	56. 151	44.960	1.00 20.02	A	C
ATOM	4289	CG	ASP	556 556	42.649	55. 255	44. 574	1.00 19.76	A	C
ATOM	4290		ASP	556	42. 723	54. 132	45. 115	1.00 19.65	A	0
ATOM	4291		ASP	556	43.470	55.666	43. 723	1.00 21.90	A	0
ATOM	4292	C	ASP	556	40.478	54. 603	46.614	1.00 20.18	A	C
ATOM ATOM	4293	0 N	ASP	556	40.856	53. 874	47. 523	1.00 19.93	A	0
ATOM	4294 4295	N CA	THR THR	557 557	39. 542	54. 246	45. 736	1.00 20.55	A	N
ATOM	4296	CB	THR	557 557	38. 835 37. 331	52. 965 53. 154	45. 820	1.00 22.31 1.00 21.37	A	C
ATOM	4297		THR	557	37. 130		45. 578		A	C
ATOM	4298		THR	557	36. 754	54. 201	46. 523	1.00 21.50	A	0
ATOM	4299	C	THR	557	39. 294		40. 323	1.00 21.28 1.00 23.72	A	C
ATOM	4300	Ŏ	THR	557	38. 606	50. 891	44. 633	1.00 25.72	A	C
ATOM	4301	N	VAL	558	40. 441	52. 105	44. 194	1.00 23.32	A A	O N
ATOM	4302	CA	VAL	558	40. 931	51. 143	43. 219	1.00 22.53	A	C
ATOM	4303	CB	VAL	558	41.970		42. 294	1.00 22.67	A	Č
ATOM	4304		VAL	558	42.540		41. 323	1.00 19.20	A	Č
ATOM	4305		VAL	558	41.323		41.547	1.00 13.20	A	Č
ATOM	4306	C	VAL	558	41.544		43. 871	1.00 21.12	A	Č
ATOM	4307	ŏ	VAL	558	42. 246	50.005	44. 871	1.00 23.71	Ÿ.	0
ATOM	4308	Ň	PHE	559	41. 261	48. 734	43. 312	1.00 25.05	A	Ň
ATOM	4309	CA	PHE	559	41.815	47. 492	43. 841	1.00 25.45	A	Č
ATOM	4310	CB	PHE	559	40. 855	46. 326	43. 584	1.00 24.60	A	Č
ATOM	4311	ĊĞ	PHE	559	41.476	44. 977	43. 808	1.00 24.75	A	Č
				-	-1. 1.0			MA- +U	* *	v

					FΙ	G. 4	- 89			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4313 4314 4315 4316 4317 4318 4319 4320 4321 4322 4323 4324 4325 4329 4330 4331 4332 4333 4334 4335 4336 4337 4338 4339 4340 4341 4342 4343 4344 4345 4346 4347 4348 4349 4340 4341 4342 4343 4344 4345 4346 4347 4348 4349 4340 4341 4342 4343 4344 4345 4346 4347 4348 4349 4340 4341 4342 4343 4344 4345 4346 4347 4348 4349 4340 4341 4342 4343 4344 4345 4346 4347 4348 4349 4340 4341 4342 4343 4344 4345 4346 4347 4348 4349 4340 4341 4342 4343 4343 4344 4345 4346 4347 4348 4349 4350 436 437 438 438 438 438 438 438 438 438	CD2 CCE2 CCCC CCNH2 CNH2 CNH2 CCCCCC CCCCC CCCCC CCCCCC CCCCCCCC CCCC	ASN ASN ASN TRP TRP TRP TRP TRP TRP	560 560 560 560 560 560 561 561 561 561 562 562 562 562 562 563 563 563 563 563 563 563 563	42. 192 41. 382 42. 810 41. 995 42. 709 43. 158 43. 250 44. 188 45. 508 46. 269 46. 885 46. 269 45. 637 45. 543 46. 112 47. 111 47. 968 46. 283 46. 139 46. 045 49. 380 49. 894 49. 999 51. 335 51. 197 50. 364 49. 881 50. 195 52. 291 52. 055 53. 375 54. 366 55. 538 56. 741 57. 367	44. 352 43. 118 43. 125 47. 246 46. 912 46. 912 46. 912 46. 912 47. 892 49. 140 50. 285 51. 536 52. 391 52. 149 53. 468 44. 255 44. 152 44. 152 44. 822 45. 916 40. 749 41. 460 42. 523 41. 916 42. 523 43. 740 44. 255 44. 152 45. 081 46. 912 47. 892 47. 491 48. 610 46. 729 47. 491 48. 537 46. 537 46. 537 46. 537 46. 537 46. 545 46. 545 47. 200 48. 575 45. 041	42. 799 45. 044 43. 021 45. 276 44. 266 43. 170 41. 943 43. 962 43. 390 42. 869 43. 310 42. 515 41. 218 43. 022 43. 980 45. 145 43. 136 43. 511 42. 635 43. 136 43. 511 42. 635 44. 203 45. 145 42. 773 41. 803 44. 274 44. 142 44. 028 45. 148 45. 054 46. 223 46. 056 46. 423 47. 178 48. 602 48. 198 47. 361	1. 00 25. 27 1. 00 28. 04 1. 00 24. 71 1. 00 26. 38 1. 00 26. 14 1. 00 27. 21 1. 00 23. 52 1. 00 20. 68 1. 00 19. 21 1. 00 20. 38 1. 00 20. 51 1. 00 20. 51 1. 00 24. 37 1. 00 24. 37 1. 00 24. 84 1. 00 23. 62 1. 00 20. 95 1. 00 18. 87 1. 00 20. 60 1. 00 19. 75 1. 00 17. 53 1. 00 20. 00 1. 00 20. 19 1. 00 18. 87 1. 00 18. 87 1. 00 18. 39 1. 00 18. 39 1. 00 18. 39 1. 00 18. 48 1. 00 19. 79 1. 00 17. 98 1. 00 17. 98 1. 00 17. 62 1. 00 17. 62 1. 00 15. 76 1. 00 13. 80 1. 00 13. 46 1. 00 12. 65	A A A A A A A A A A A A A A A A A A A	CCCCCONCCCNCNNCONCCCCCONCCCONCONCCCCCCCC
ATOM ATOM	4355 C	E1 1 Z2 1	TRP	563 563	58. 440 59. 439	45. 189 47. 128	48. 217 49. 453	1.00 11.34 1.00 14.40	A A	N C
ATOM ATOM ATOM			TRP TRP TRP	563 563 563	58. 252 59. 291	49. 204 48. 476	49. 046 49. 664	1.00 16.29 1.00 14.18	A A	C
ATOM ATOM	4359 O 4360 N	1	TRP NLA	563 564	53. 728 54. 048 52. 813	45. 672 44. 910 46. 620		1. 00 17. 48 1. 00 18. 93 1. 00 16. 80	A A A	C O N
									••	**

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(Continued) FIG. 4-90 46.838 49.232 C **ATOM** 4361 CA ALA 564 52.151 1.00 17.11 C **ATOM** 4362 CB ALA 564 51.248 48.068 49.153 1.00 16.72 A 4363 C **ALA** 51.341 45.616 49.655 1.00 17.89 A C ATOM 564 45.256 0 4364 0 ALA 564 51.322 50.834 1.00 15.94 ATOM Α 50.676 44.983 48.691 1.00 18.77 4365 N THR 565 N ATOM Α 48.977 C 4366 CA THR 565 49.870 43.801 1.00 19.59 A **ATOM** C 4367 CB THR 565 49.368 43.131 47.689 1.00 20.01 A ATOM 4368 OG1 THR 48.606 44.069 46.922 1.00 19.76 0 **ATOM** 565 A 4369 CG2 THR 48.496 41.922 48.027 1.00 19.34 C **ATOM** 565 Α C 4370 C 50.718 42.793 49.739 1.00 21.27 **ATOM** THR 565 A 4371 42.252 50.760 1.00 22.29 0 ATOM 0 565 50.290 THR Α 4372 51.924 42.548 49.234 1.00 22.25 **ATOM** N TYR 566 A N 4373 41.615 1.00 23.40 52.848 49.864 **ATOM** CA TYR 566 C Α 54.029 41.324 48.923 $\begin{array}{c} C \\ C \\ C \\ C \end{array}$ **ATOM** 4374 CB TYR 566 1.00 25.18 A 1.00 25.40 4375 CG TYR 566 55.369 41.218 49.616 **ATOM** A 4376 56.297 42.262 49.547 1.00 25.62 **ATOM** CD1 TYR 566 Α 50.226 4377 CE1 **TYR** 57.513 42.196 1.00 26.85 **ATOM** 566 Α 55.690 50.382 C **ATOM** 4378 CD2 TYR 566 40.101 1.00 26.99 A Ċ 4379 CE2 TYR 566 56.903 40.023 51.073 1.00 29.74 ATOM Α 4380 57.809 41.074 50.991 ATOM CZ TYR 566 1.00 30.16 A C 4381 0HTYR 58.997 **ATOM** 566 40.998 51.688 1.00 32.61 A 0 4382 C TYR 566 53.369 51.212 ATOM 42.116 1.00 23.06 C Α 53.458 4383 0 566 41.350 52.170 1.00 21.96 **ATOM** TYR Α 0 4384 N LEU 53.716 43.396 51.288 **ATOM** 567 1.00 23.28 N Α 52.532 4385 54.237 C **ATOM** CA LEU 567 43.949 1.00 24.50 A ATOM 4386 CB LEU 54.588 52.359 567 45.429 1.00 22.74 C A CG LEU Č **ATOM** 4387 567 55.717 45.769 51.378 1.00 23.15 A **ATOM** 4388 CD1 LEU 55.833 47.279 51.263 C 567 1.00 20.37 A 4389 CD2 LEU 567 57.038 45.158 51.850 C **ATOM** 1.00 21.42 A 4390 C 53.243 1.00 26.32 **ATOM** LEU 567 43.786 53.675 Α C **ATOM** 4391 0 LEU 567 53.635 43.595 54.824 1.00 27.44 0 A **ATOM** 4392 51.955 N ALA 568 43.857 53.361 1.00 26.96 A N 4393 50.930 1.00 27.44 ATOM CA ALA 568 43.712 54.383 C A **ATOM** 4394 CB ALA 568 49.684 44.481 53.984 1.00 26.54 C Α 4395 C 568 50.584 42.242 1.00 29.12 ATOM ALA 54.606 A C 4396 **ATOM** 0 ALA 568 50.483 41.782 55.748 1.00 28.80 Α 0 4397 N ATOM SER 569 50.417 41.506 53.509 1.00 28.58 A N 4398 569 **ATOM** CA SER 50.062 40.094 53.586 1.00 28.31 C A **ATOM** 4399 CB SER 569 49.750 39.553 52.191 1.00 28.85 C Α 4400 0G 38.174 52.247 1.00 30.69 **ATOM** SER 569 49.420 Α 0 **ATOM** 4401 C SER 569 51.110 39.204 54.236 1.00 27.43 C A 0 38.427 **ATOM** 4402 SER 569 50.800 55.133 1.00 28.44 Α 0 4403 N THR 570 52.350 39.311 53.781 1.00 27.24 **ATOM** N 38.483 4404 CA 570 53.420 54.314 1.00 27.02 ATOM THR C 570 54.410 38.094 53.199 4405 CB THR 1.00 26.90 C **ATOM** A 52.248 **ATOM** 4406 OG1 THR 570 53.749 37.250 1.00 27.63 A 0 **ATOM** CG2 THR 570 55.611 37.369 53.774 1.00 23.88 C 4407 A 1.00 27.34 54.203 C C THR 570 39.110 55.459 Α ATOM 4408

SUBSTITUTE SHEET (RULE 26)

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54.362

4409

ATOM

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THR

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	,	FIG. 4-91	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4411 CA GLU 571 4412 CB GLU 571 4413 CG GLU 571 4414 CD GLU 571 4415 OE1 GLU 571 4416 OE2 GLU 571 4417 C GLU 571 4418 O GLU 571 4419 N ASN 572 4420 CA ASN 572 4421 CB ASN 572 4422 CG ASN 572 4423 OD1 ASN 572 4424 ND2 ASN 572 4425 C ASN 572 4426 O ASN 572 4427 N ILE 573 4428 CA ILE 573 4429 CB ILE 573	54. 686 40. 329 55. 253 1. 00 26. 71 55. 480 41. 020 56. 259 1. 00 25. 23 56. 402 42. 040 55. 583 1. 00 24. 64 57. 287 41. 472 54. 473 1. 00 25. 43 58. 238 40. 392 54. 966 1. 00 27. 45 58. 582 40. 421 56. 164 1. 00 28. 11 58. 656 39. 527 54. 158 1. 00 27. 18 54. 643 41. 715 57. 329 1. 00 24. 50 55. 188 42. 368 58. 213 1. 00 24. 29 53. 324 41. 576 57. 247 1. 00 24. 39 52. 425 42. 191 58. 223 1. 00 24. 96 52. 557 41. 486 59. 569 1. 00 25. 44 52. 139 40. 033 59. 507 1. 00 29. 03 52. 711 39. 187 60. 192 1. 00 30. 88 51. 128 39. 734 58. 694 1. 00 29. 67 52. 683 43. 681 58. 419 1. 00 25. 32 52. 642 44. 178 59. 545 1. 00 25. 48 53. 208 45. 824 <	A N A C A C A C A C A C A C A C A C A C A C
ATOM ATOM	4430 CG2 ILE 573 4431 CG1 ILE 573	54.715 47.669 56.584 1.00 22.90	A C
ATOM	4432 CD1 ILE 573	55. 622 45. 365 56. 800 1. 00 25. 08 56. 805 45. 636 55. 900 1. 00 25. 36	A C A C
ATOM	4433 C ILE 573	51. 992 46. 621 56. 875 1. 00 25. 22	A. C
ATOM	4434 O ILE 573	51. 353 46. 249 55. 891 1. 00 24. 86	A O
ATOM ATOM	4435 N ILE 574 4436 CA ILE 574	51.681 47.718 57.557 1.00 24.59	A N
ATOM	4436 CA ILE 574 4437 CB ILE 574	50. 557 48. 555 57. 159 1. 00 26. 14	A C
ATOM	4438 CG2 ILE 574	49. 926 49. 297 58. 359 1. 00 25. 88 48. 798 50. 190 57. 874 1. 00 26. 06	A C
ATOM	4439 CG1 ILE 574	48. 798 50. 190 57. 874 1. 00 26. 06 49. 399 48. 304 59. 386 1. 00 27. 36	A C
ATOM	4440 CD1 ILE 574	48.794 48.968 60.607 1.00 29.19	A C
ATOM	4441 C ILE 574	51.064 49.619 56.191 1.00 27.12	A C A C
ATOM	4442 O ILE 574	51. 799 50. 524 56. 591 1. 00 28. 97	A Ö
ATOM ATOM	4443 N VAL 575 4444 CA VAL 575	50. 683 49. 521 54. 924 1. 00 25. 92	A N
ATOM	4444 CA VAL 575 4445 CB VAL 575	51. 128 50. 517 53. 962 1. 00 24. 87	A C
ATOM	4446 CG1 VAL 575	51. 387 49. 904 52. 569 1. 00 24. 76 51. 973 50. 966 51. 644 1. 00 20. 17	A C
ATOM	4447 CG2 VAL 575	51. 973 50. 966 51. 644 1. 00 20. 17 52. 320 48. 707 52. 690 1. 00 22. 12	A C
ATOM	4448 C VAL 575	50.054 51.585 53.837 1.00 25.21	A C A C
ATOM	4449 0 VAL 575	48. 929 51. 312 53. 405 1. 00 25. 63	A 0
ATOM ATOM	4450 N ALA 576	50. 403 52. 804 54. 216 1. 00 23. 75	A N
ATOM	4451 CA ALA 576 4452 CB ALA 576	49. 456 53. 893 54. 152 1. 00 23. 56	ACC
ATOM	4453 C ALA 576	49. 255 54. 477 55. 540 1. 00 23. 43 49. 879 54. 988 53. 180 1. 00 24. 06	A C
ATOM	4454 0 ALA 576	49. 879 54. 988 53. 180 1. 00 24. 06 51. 056 55. 139 52. 860 1. 00 22. 16	A C
ATOM	4455 N SER 577	48. 888 55. 740 52. 710 1. 00 24. 49	A O A N
ATOM	4456 CA SER 577	49. 095 56. 852 51. 796 1. 00 23. 11	A C
ATOM	4457 CB SER 577	48. 793 56. 428 50. 362 1. 00 23. 06	A C
ATOM	4458 OG SER 577	49. 750 55. 475 49. 921 1. 00 22. 88	A 0

-					FI	G. 4	- 92			(Continued)
ATOM ATOM	4459 4460	C 0	SER SER	577 577	48. 149 47. 075	57. 947 57. 662	52. 248 52. 768	1.00 22.90 1.00 24.22	A A	C 0
ATOM	4461	Ň	PHE	578	48. 546	59.196	52.046	1.00 23.49	A	Ň
ATOM	4462	CA	PHE	578	47.748	60.337	52.479	1.00 21.77	Ā	C
ATOM	4463	CB	PHE	578	48. 313	60.829	53.804	1.00 21.41	Α	C
ATOM	4464	CG	PHE	578	47. 585	62.005	54. 383	1.00 22.79	Α	C .
ATOM	4465		PHE	578	46. 429	61.820	55. 144	1.00 20.60	A	C
ATOM	4466		PHE	578	48. 080	63. 291	54. 209	1.00 19.79	A	C
ATOM	4467		PHE	578 579	45. 783	62.901	55. 730	1.00 21.26	A	C
ATOM ATOM	4468 4469	CZ	PHE PHE	578 578	47. 441 46. 288	64. 381 64. 186	54.790 55.556	1.00 20.94 1.00 20.70	A	C C C C
ATOM	4470	C	PHE	578	47. 723	61.502	51.480	1.00 20.70	A A	C
ATOM	4471	Õ	PHE	578	48. 766	61.909	50. 973	1.00 21.14	A	Ö
ATOM	4472	Ň	ASP	579	46. 533	62.041	51. 212	1.00 19.89	Ä	Ň
ATOM	4473	CA	ASP	579	46. 389	63.173	50.302	1.00 18.01	Ä	Ĉ
ATOM	4474	CB	ASP	579	45. 191	62.985	49.371	1.00 17.01	Α	C C C
ATOM	4475	CG	ASP	579	45. 334	61.777	48.455	1.00 21.86	Α	C
ATOM	4476		ASP	579	46. 424	61.583	47.873	1.00 22.87	Α	0
ATOM	4477		ASP	579	44. 342	61.024	48. 299	1.00 23.17	A	0
ATOM	4478	C	ASP	579	46. 211	64. 474	51.092	1.00 18.10	A	C
ATOM ATOM	4479 4480	O N	ASP GLY	579 580	45. 103	64. 823 65. 189	51.493	1.00 20.42	A	0
ATOM	4481	CA	GLY	580	47. 306 47. 238	66. 439	51. 313 52. 044	1.00 17.22 1.00 15.14	A A	N
ATOM	4482	C	GLY	580	47. 065	67.610	51. 098	1.00 15.14	A	C C
ATOM	4483	ŏ	GLY	580	46. 544	67.462	49. 993	1.00 17.18	A	Õ
ATOM	4484	N	ARG	581	47. 495	68.786	51. 528	1.00 15.90	Ä	N
ATOM	4485	CA	ARG	581	47. 377	69.970	50.701	1.00 15.52	Ā	C
ATOM	4486	CB	ARG	581	47. 956	71.172	51.444	1.00 16.17	Α	C
ATOM	4487	CG	ARG	581	47. 072	71.645	52.585	1.00 16.05	Α	C
ATOM	4488	. CD	ARG	581	47. 756	72.653	53. 467	1.00 14.87	A	C
ATOM	4489	NE CZ	ARG	581	48.617	71.990	54. 441	1.00 18.25	A	N
ATOM ATOM	4490	CZ	ARG	581	49. 321	72.624	55. 375	1.00 19.44	A	C
ATOM	4491 4492		ARG ARG	581 581	49. 268 50. 075	73. 952 71. 933	55. 463 56. 224	1.00 20.41 1.00 15.76	A	N
ATOM	4493	C	ARG	581	48. 107	69. 742	49. 386	1.00 15.76	A A	N C
ATOM	4494	ŏ	ARG	581	49. 193	69. 158	49. 357	1.00 17.49	Ä	Õ
ATOM	4495	Ň	GLY	582	47. 495	70. 192	48. 295	1.00 18.96	A	Ň
ATOM	4496	CA	GLY	582	48.094	70.022	46. 987	1.00 17.63	Ä	. Ĉ
ATOM	4497	C	GLY	582	47. 511	68.842	46. 231	1.00 18.54	Α	C ·
MOTA	4498	0	GLY	582	47.673	68.757	45.017	1.00 18.99	Α	0
ATOM	4499	N	SER	583	46.842	67.923	46.925	1.00 18.00	Α	N
ATOM	4500	CA	SER	583	46. 258	66. 765	46. 247	1.00 18.46	A	C
ATOM	4501	CB	SER	583	45.842	65.700	47. 269	1.00 18.34	A	C
ATOM	4502 4503	OG C	SER SER	583 583	45.058	66. 253	48. 303	1.00 19.12	A.	0
ATOM ATOM	4503 4504	0	SER	583	45. 068 44. 601	67. 218 68. 344	45. 392 45. 536	1.00 18.03 1.00 17.42	A A	C
ATOM	4505	N	GLY	584	44. 570	66. 355	45. 536	1.00 17.42	A A	O N
ATOM	4506	CA	GLY	584	43. 481	66.779	43. 637	1.00 17.04	A	C
ATOM	4507	C	GLY	584	42. 052	66. 293	43. 827	1.00 19.49	A	č
									••	-

	FIG. 4-93	(Continued)
ATOM 4508 O GLY 584 ATOM 4509 N TYR 585 ATOM 4510 CA TYR 585 ATOM 4511 CB TYR 585 ATOM 4512 CG TYR 585 ATOM 4513 CD1 TYR 585 ATOM 4514 CE1 TYR 585 ATOM 4515 CD2 TYR 585 ATOM 4516 CE2 TYR 585 ATOM 4517 CZ TYR 585 ATOM 4518 OH TYR 585 ATOM 4519 C TYR 585 ATOM 4520 O TYR 585 ATOM 4521 N GLN 586 ATOM 4522 CA GLN 586 ATOM 4522 CA GLN 586 ATOM 4523 CB GLN 586 ATOM 4524 CG GLN 586 ATOM 4525 CD GLN 586 ATOM 4526 OE1 GLN 586 ATOM 4527 NE2 GLN 586 ATOM 4528 C GLN 586 ATOM 4529 O GLN 586 ATOM 4520 O GLN 586 ATOM 4521 N GLY 587 ATOM 4531 CA GLY 587 ATOM 4530 N GLY 587 ATOM 4531 CA GLY 587 ATOM 4531 CA GLY 587 ATOM 4532 C GLY 587 ATOM 4533 O GLY 587 ATOM 4534 N ASP 588 ATOM 4535 CA ASP 588 ATOM 4536 CB ASP 588 ATOM 4537 CG ASP 588 ATOM 4538 OD1 ASP 588 ATOM 4539 OD2 ASP 588 ATOM 4530 N GLY 587 ATOM 4531 CA GLY 587 ATOM 4534 N ASP 588 ATOM 4535 CA ASP 588 ATOM 4536 CB ASP 588 ATOM 4537 CG ASP 588 ATOM 4538 OD1 ASP 588 ATOM 4539 OD2 ASP 588 ATOM 4540 C ASP 588 ATOM 4540 C ASP 588 ATOM 4541 O ASP 588 ATOM 4542 N LYS 589 ATOM 4545 CG LYS 589 ATOM 4546 CD LYS 589 ATOM 4547 CE LYS 589 ATOM 4548 NZ LYS 589 ATOM 4549 C LYS 589 ATOM 4540 C LYS 589 ATOM 4541 N LE 590 ATOM 4555 CGI ILE 590 ATOM 4555 CGI ILE 590 ATOM 4555 CGI ILE 590 ATOM 4556 CDI ILE 590 ATOM 4556 CDI ILE 590	## F I G. 4 - 9 3 ## 1.724 65. 570 44. 767 1. 00 21. 57 ## 1.91 66. 735 42. 917 1. 00 19. 76 ## 39. 782 66. 362 42. 906 1. 00 18. 53 ## 39. 673 64. 859 42. 663 1. 00 18. 57 ## 40. 578 64. 401 41. 550 1. 00 18. 83 ## 40. 439 64. 914 40. 260 1. 00 19. 48 ## 41. 300 64. 533 39. 235 1. 00 18. 11 ## 41. 606 63. 490 41. 789 1. 00 19. 81 ## 42. 476 63. 100 40. 769 1. 00 17. 71 ## 42. 313 63. 626 39. 497 1. 00 18. 76 ## 43. 150 63. 232 38. 481 1. 00 20. 70 ## 38. 997 66. 751 44. 152 1. 00 18. 81 ## 38. 046 66. 067 44. 521 1. 00 17. 85 ## 39. 382 67. 861 44. 783 1. 00 20. 25 ## 38. 708 68. 345 45. 986 1. 00 20. 04 ## 39. 475 66. 412 47. 279 1. 00 20. 60 ## 40. 481 66. 095 48. 363 1. 00 24. 77 ## 40. 441 66. 029 49. 548 1. 00 23. 60 ## 42. 044 65. 919 47. 962 1. 00 25. 12 ## 38. 619 69. 869 46. 024 1. 00 22. 06 ## 43. 424 70. 455 47. 092 1. 00 23. 83 ## 37 70. 518 44. 877 1. 00 21. 79 ## 40. 073 72. 623 44. 883 1. 00 21. 24 ## 40. 073 72. 623 44. 883 1. 00 21. 24 ## 40. 465 76. 213 41. 211 1. 00 25. 77 ## 41. 157 74. 155 44. 397 1. 00 22. 09 ## 41. 287 75. 763 43. 382 1. 00 22. 09 ## 41. 287 75. 663 44. 397 1. 00 21. 25 ## 41. 157 74. 155 41. 599 1. 00 24. 41 ## 41. 955 75. 079 45. 675 1. 00 23. 77 ## 41. 157 74. 155 41. 599 1. 00 24. 41 ## 41. 955 75. 079 45. 675 1. 00 23. 77 ## 41. 157 74. 155 41. 599 1. 00 24. 41 ## 41. 955 75. 079 45. 675 1. 00 23. 77 ## 41. 157 74. 155 41. 599 1. 00 24. 29 ## 42. 494 75. 386 46. 716 1. 00 22. 97 ## 40. 494 76. 173 50. 365 1. 00 22. 97 ## 40. 497 76. 638 52. 784 1. 00 36. 09 ## 42. 494 78. 326 48. 881 1. 00 36. 83 ## 42. 705 74. 611 48. 468 1. 00 22. 93 ## 43. 710 75. 086 46. 6716 1. 00 23. 77 ## 41. 905 75. 616 49. 037 1. 00 24. 26 ## 42. 494 78. 326 48. 881 1. 00 36. 83 ## 42. 705 74. 611 48. 468 1. 00 22. 97 ## 40. 497 75. 648 48. 607 1. 00 21. 76 ## 41. 901 71. 025 50. 126 1. 00 21. 94 ## 41. 901 71. 025 50. 126 1. 00 21. 94 ## 41. 901 71. 025 50. 126 1. 00 21. 94 ## 41. 200 69. 720 50. 478 1. 00 22. 25	Continued) A O O O O O O O O O O O O O O O O O O
1001	41. 901 71. 025 50. 126 1. 00 21. 94	A C

										(Con	tinued)
					FIC	3. 4	- 94				
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4557 4558 4559 4560 4561 4562 4563 4564 4565 4566 4567 4570 4571 4572 4573 4574 4577 4578	ND1 CE1	ILE ILE MET MET MET MET HIS HIS HIS HIS HIS ALA ALA	590 590 591 591 591 591 591 592 592 592 592 592 592 593	44. 537 45. 711 44. 157 45. 127 44. 406 45. 309 44. 403 44. 237 46. 112 47. 289 45. 636 46. 502 45. 713 45. 296 45. 604 44. 471 44. 289 44. 965 47. 197 47. 842 47. 076	72. 093 71. 960 72. 071 71. 846 71. 567 71. 000 70. 746 72. 436 72. 997 72. 771 74. 228 75. 386 76. 560 76. 361 75. 390 77. 243 76. 825 75. 703 75. 817 76. 865 75. 012 75. 349	47. 562 47. 901 46. 291 45. 232 43. 917 42. 838 41. 309 40. 732 45. 051 44. 791 45. 200 45. 035 44. 455 43. 032 42. 139 42. 368 41. 128 40. 962 46. 319 46. 362 47. 367 48. 628	1. 00 22. 32 1. 00 23. 51 1. 00 21. 59 1. 00 21. 80 1. 00 21. 85 1. 00 22. 76 1. 00 22. 84 1. 00 21. 43 1. 00 21. 21 1. 00 21. 21 1. 00 21. 21 1. 00 22. 32 1. 00 24. 65 1. 00 25. 75 1. 00 25. 75 1. 00 25. 78 1. 00 21. 38 1. 00 20. 84 1. 00 20. 43	A A A A A A A A A A A A A A A A A A A	CONCCCSCCONCCCCNCNCON	atinued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4579 4580 4581 4582 4583 4584 4585 4586 4587 4590 4591 4592 4593 4594 4595 4596 4597 4598	CB CON CACB CG2 CG1 CD1 CON CACB CGOD1 ND2 CON	ALA ALA ILE ILE ILE ILE ILE ILE ILE ASN ASN ASN ASN ASN ASN ASN ASN	593 593 594 594 594 594 594 595 595 595 595 595	47. 360 49. 241 49. 940 49. 736 51. 176 51. 617 51. 467 50. 814 50. 951 51. 658 52. 849 50. 746 51. 119 49. 977 50. 300 50. 640 50. 191 52. 395 52. 442 53. 421	74. 349 75. 361 76. 126 74. 522 74. 446 73. 021 72. 051 72. 581 71. 106 75. 410 75. 434 76. 200 77. 137 78. 114 79. 072 78. 652 80. 364 77. 921 78. 688 77. 715	49. 710 48. 427 49. 081 47. 518 47. 248 46. 816 47. 966 45. 590 45. 243 46. 169 45. 854 45. 606 44. 547 44. 265 43. 128 42. 024 43. 394 44. 860 45. 824 44. 031	1. 00 18. 24 1. 00 19. 92 1. 00 21. 91 1. 00 19. 47 1. 00 20. 49 1. 00 19. 36 1. 00 19. 38 1. 00 21. 33 1. 00 22. 55 1. 00 19. 88 1. 00 17. 79 1. 00 20. 03 1. 00 21. 76 1. 00 20. 68 1. 00 21. 80 1. 00 22. 78 1. 00 22. 74 1. 00 22. 44 1. 00 22. 52	A A A A A A A A A A A A A A A A A A A	C C C O N C C C C O N C C O N C O N C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4599 4600 4601 4602 4603 4604 4605	CA CB CG CD NE CZ NH1	ARG ARG ARG ARG ARG ARG ARG	596 596 596 596 596 596	54. 550 53. 894 53. 398 54. 479 55. 467	78. 378 79. 898 80. 426 81. 856 82. 760 83. 112 82. 635	44. 171 44. 141 42. 880 43. 096 43. 482 42. 671 41. 431	1. 00 22. 41 1. 00 21. 28 1. 00 21. 31 1. 00 22. 01 1. 00 20. 88 1. 00 21. 35 1. 00 22. 62	A A A A A A	C C C N C N	

					FIG. 4-95	(Continued)
ATOM	4606		I2 ARG	596	56. 427 83. 924 43. 096 1. 00 19. 92 A	N
ATOM	4607		ARG	596	55. 492 77. 982 45. 440 1. 00 21. 53 A	C
ATOM ATOM	4608		ARG	596	56. 482 78. 611 45. 804 1. 00 20. 59 A	0
ATOM	4609 4610		ARG	597	55. 046 76. 930 46. 107 1. 00 21. 66 A	N
ATOM	4611			597	55. 705 76. 512 47. 331 1. 00 21. 98 A	C
ATOM	4612			597 597	54. 943 77. 061 48. 539 1. 00 23. 55 A	Ç.
ATOM	4613			597	55.184 78.547 48.776 1.00 28.20 A	C
ATOM	4614			597	56.611 78.813 49.264 1.00 30.86 A 56.891 80.239 49.414 1.00 34.81 A	C
ATOM	4615			597	FR 081 04 000 10 10	N
ATOM	4616		1 ARG	597	F7 011 00 000 (F 110	C
ATOM	4617		2 ARG	597	F7 886 00 00F 10 0F0	N
ATOM	4618		ARG	597	EE 000 EE 014 IE 150	N
ATOM	4619		ARG	597	55. 523 74. 423 48. 487 1. 00 20. 79 A	C
ATOM	4620	N	LEU	598	56. 400 74. 398 46. 404 1. 00 19. 44 A	O N
ATOM	4621	CA		598	56. 649 72. 963 46. 387 1. 00 18. 48 A	C
ATOM	4622	CB	LEU	598	57. 142 72. 545 45. 003 1. 00 18. 20 A	Č
ATOM	4623	CG		598	56.119 72.007 43.994 1.00 19.27 A	č
ATOM `	4624		LEU	598	54. 800 72. 731 44. 107 1. 00 19. 49 A	č
ATOM .	4625		2 LEU	598	56.691 72.135 42.595 1.00 18.24 A	č
ATOM	4626	C	LEU	598	57.692 72.617 47.450 1.00 19.10 A	č
ATOM	4627	0	LEU	598	58. 644 73. 363 47. 679 1. 00 19. 27 A	Ö
ATOM	4628	N	GLY	599	57. 506 71. 485 48. 108 1. 00 19. 24 A	N
ATOM ATOM	4629 4630	CA	GLY	599	58. 440 71. 090 49. 138 1. 00 20. 34 A	C
ATOM	4631	C	GLY	599	58. 055 71. 622 50. 508 1. 00 21. 76 A	C
ATOM	4632	O N	GLY THR	599	58. 882 71. 640 51. 422 1. 00 23. 58 A	0
ATOM	4633	CA	THR	600	56. 811 72. 061 50. 666 1. 00 21. 02 A	N
ATOM	4634	CB	THR	600 600	56. 381 72. 578 51. 958 1. 00 21. 20 A	C
ATOM	4635	0G1		600	56. 039 74. 082 51. 874 1. 00 21. 28 A	C
ATOM	4636		THR	600	54. 887 74. 271 51. 052 1. 00 25. 68 A	0
ATOM	4637	C	THR	600	57. 192 74. 856 51. 264 1. 00 21. 23 A 55. 201 71. 810 52. 557 1. 00 21. 38 A	C
ATOM	4638	ŏ ·	THR	600	EE 200 70 704 E2 400 4 00 22 1-	C
ATOM	4639	N	PHE	601	E9 009 70 0F0 F0 440 4 00 04 15	0
ATOM	4640			601	E9 900 71 701 E9 000 1 00 00 00	N
ATOM	4641	CB	PHE	601	E1 E40 70 400 F0 040 4 00 04 00	C
ATOM	4642	CG	PHE	601	51. 556 73. 935 53. 077 1. 00 24. 93 A	C C
ATOM	4643		PHE	601	51.052 74.923 52.236 1.00 28.07 A	C
ATOM	4644		PHE	601	52.105 74.308 54.299 1.00 26.83 A	C
ATOM	4645		PHE	601	51. 100 76. 271 52. 603 1. 00 29. 10 A	Č
ATOM	4646		PHE	601	52. 160 75. 650 54. 680 1. 00 28. 02 A	č
ATOM	4647	CZ	PHE	601	51.658 76.636 53.830 1.00 28.61 A	č
ATOM	4648	C	PHE	601	52. 623 70. 265 52. 635 1. 00 22. 45 A	č·.
ATOM	4649	0	PHE	601	52. 235 69. 451 53. 470 1. 00 22. 89 A	0.
ATOM	4650	N	GLU	602	52.884 69.931 51.374 1.00 22.76 A	N
ATOM	4651	CA	GLU	602	52. 712 68. 556 50. 931 1. 00 21. 82 A	Ĉ
ATOM	4652	CB	GLU	602	52.956 68.418 49.422 1.00 22.43 A	C
ATOM	4653	CC	GLU	602	54. 396 68. 559 48. 974 1. 00 27. 44 A	C
ATOM	4654	CD	GLU	602	54.872 70.002 48.893 1.00 29.71 A	C

				FIG. 4-96	(Continued)
ATOM	4655	OE1 GLU	602	FA 751 70 749 40 001 1 00 01 77	0
ATOM	4656		602	54. 751 70. 743 49. 891 1. 00 31. 66 A	0
ATOM	4657		602	55. 379 70. 392 47. 822 1. 00 31. 46 A	0
ATOM	4658		602	53. 663 67. 657 51. 698 1. 00 21. 67 A	C
ATOM	4659		603	53. 386 66. 473 51. 899 1. 00 22. 33 A	0
ATOM	4660		603	54. 777 68. 229 52. 146 1. 00 20. 78 A 55. 772 67. 468 52. 897 1. 00 20. 76 A	N
ATOM	4661	CB VAL	603	F7 1F0 40 100 FD 000 1 00 10	C
ATOM	4662	CG1 VAL	603	EO 105 05 05 50 010 1 00 15	C
ATOM	4663	CG2 VAL	603	F. 000 00 100 F. 00F	C
ATOM	4664	C VAL	603	EE 000 00 000 E4 004 4 00 54	C
ATOM	4665	0 VAL	603	FF 979 00 00F F1 010 1 00 F2	C
ATOM	4666	N GLU	604	FF 000 00 101 F1 0F1	0 N
ATOM	4667	CA GLU	604	55. 009 68. 481 54. 951 1. 00 24. 70 A 54. 594 68. 518 56. 341 1. 00 27. 84 A	N C
ATOM	4668	CB GLU	604	54. 322 69. 964 56. 770 1. 00 30. 83 A	C C
ATOM	4669	CG GLU	604	55. 572 70. 808 56. 924 1. 00 37. 92 A	C
ATOM	4670	CD GLU	604	56. 449 70. 355 58. 091 1. 00 43. 63 A	C
ATOM	4671	OE1 GLU	604	57. 505 70. 989 58. 328 1. 00 46. 30 A	Õ
ATOM	4672	OE2 GLU	604	56. 083 69. 368 58. 773 1. 00 45. 85 A	Ö
ATOM	4673	C GLU	604	53. 349 67. 669 56. 553 1. 00 27. 28 A	č
ATOM	4674	0 GLU	604	53. 270 66. 909 57. 517 1. 00 28. 68 A	ŏ
ATOM	4675	N ASP	605	52. 381 67. 786 55. 650 1. 00 25. 92 A	Ň
ATOM	4676	CA ASP	605	51.151 67.021 55.785 1.00 25.72 A	Ċ
ATOM	4677	CB ASP	605	50. 144 67. 436 54. 713 1. 00 24. 61 A	Č
ATOM	4678	CG ASP	605	49. 576 68. 832 54. 963 1. 00 23. 36 A	Č.
ATOM	4679	OD1 ASP	605	48. 677 69. 267 54. 215 1. 00 23. 15 A	0
ATOM	4680	OD2 ASP	605	50.036 69.499 55.914 1.00 21.27 A	0
ATOM	4681	C ASP	605	51. 379 65. 515 55. 783 1. 00 26. 18 A	C
ATOM ATOM	4682	O ASP	605	50. 646 64. 779 56. 439 1. 00 28. 35 A	0
ATOM	4683 4684	N GLN CA GLN	606	52. 394 65. 051 55. 063 1. 00 26. 16 A	N
ATOM	4685	CA GLN CB GLN	606	52. 704 63. 627 55. 056 1. 00 25. 29 A	C
ATOM	4686	CG GLN	606	53. 788 63. 302 54. 026 1. 00 24. 18 A	C
ATOM	4687	CO GLN	606 606	53. 305 63. 332 52. 596 1. 00 24. 92 A	C
ATOM	4688	OE1 GLN	606	52. 206 62. 321 52. 330 1. 00 24. 81 A 52. 373 61. 122 52. 560 1. 00 25. 31 A	C
ATOM	4689	NE2 GLN	606	51 075 69 901 F1 940 1 00 9F 44	0
ATOM	4690	C GLN	606	E9 907 C9 9C0 FC 447 1 00 05 15	N
ATOM	4691	O GLN	606	E9 990 69 999 F7 999 4 99 97 17	C
ATOM	4692	N ILE	607	E4 000 04 180 07 001 1 00 00 01	0
ATOM	4693	CA ILE	607	E4 C07 C9 O1F F0 O0F 1 00 00 00	N
ATOM	4694	CB ILE	607	EE (20 CE 000 E0 E00 1 00 00 01	C
ATOM	4695	CG2 ILE	607	EC 105 CA 770 00 110 1 00 00	C
ATOM	4696	CG1 ILE	607	EC 700 C4 077 F7 004 1 00 00 00	C
ATOM	4697	CD1 ILE	607	F7 700 00 000 F7 001 1 00 00 01	C
ATOM	4698	C ILE	607	C2 470	C
ATOM	4699	0 ILE	607	53. 470 63. 963 69. 355 1. 00 29. 50 A 53. 359 63. 093 60. 226 1. 00 27. 80 A	C 0
ATOM	4700	N GLU	608	52. 619 64. 978 59. 239 1. 00 30. 32 A	N N
ATOM	4701	CA GLU	608	51.508 65.099 60.164 1.00 32.21 A	C
ATOM	4702	CB GLU	608	50. 705 66. 379 59. 919 1. 00 33. 05 A	C
ATOM	4703	CG GLU	608	49. 578 66. 581 60. 936 1. 00 34. 99 A	C
				TITLE TITLE IN	•

ATOM 4704 CD GLU 608 50.054 66.482 62.389 1.00 38.42 A C ATOM 4705 0BI GLU 608 49.197 66.454 63.302 1.00 37.67 A 0 ATOM 4706 0B2 GLU 608 51.285 66.435 62.625 1.00 40.64 A 0 ATOM 4707 C GLU 608 50.506 63.891 60.012 1.00 32.76 A C ATOM 4708 0 GLU 608 49.889 63.527 60.947 1.00 33.47 A 0 ATOM 4709 N ALA 609 50.643 63.277 58.836 1.00 31.347 A 0 ATOM 4701 CA ALA 609 49.827 62.090 58.595 1.00 30.73 A C ATOM 4711 CB ALA 609 49.827 62.090 58.595 1.00 30.73 A C ATOM 4712 C ALA 609 49.827 62.090 58.595 1.00 30.73 A C ATOM 4712 C ALA 609 50.355 60.988 59.472 1.00 30.16 A C ATOM 4711 CB ALA 609 49.883 61.682 57.123 1.00 28.50 A C ATOM 4714 N ALA 610 51.674 60.803 59.479 1.00 29.56 A N ATOM 4715 CA ALA 610 52.310 59.758 60.274 1.00 28.48 A C ATOM 4716 CB ALA 610 52.310 59.758 60.274 1.00 28.48 A C ATOM 4718 O ALA 610 51.930 59.886 61.743 1.00 27.67 A C ATOM 4719 N ARG 611 52.230 59.886 61.743 1.00 27.67 A C ATOM 4719 N ARG 611 52.230 59.886 61.743 1.00 27.67 A C ATOM 4719 N ARG 611 52.025 61.094 62.282 1.00 28.98 A C ATOM 4720 CA ARG 611 51.830 59.886 61.743 1.00 29.284 A N ATOM 4721 CB ARG 611 51.830 59.886 61.743 1.00 29.286 A C ATOM 4722 CG ARG 611 51.830 59.886 61.743 1.00 29.292 A C ATOM 4723 CD ARG 611 53.281 64.799 64.032 1.00 29.99 A C ATOM 4723 CD ARG 611 53.239 63.291 64.032 1.00 29.99 A C ATOM 4723 CD ARG 611 53.239 63.291 64.032 1.00 29.99 A C ATOM 4724 NB ARG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4724 NB ARG 611 54.980 66.384 63.378 1.00 29.99 A C ATOM 4723 CD ARG 611 54.980 66.384 63.378 1.00 29.99 A C ATOM 4724 NB ARG 611 54.980 66.384 63.378 1.00 29.99 A C ATOM 4724 NB ARG 611 54.980 66.384 63.378 1.00 29.99 A C ATOM 4725 CB MIL ARG 611 54.980 66.384 63.378 1.00 29.99 A C ATOM 4728 CB MIL ARG 611 54.980 66.384 63.378 1.00 29.99 A C ATOM 4728 CB MIL ARG 611 54.980 66.384 63.378 1.00 29.99 A C ATOM 4728 CB MIL ARG 611 54.980 66.384 63.378 1.00 29.99 A C ATOM 4735 OBI GLN 612 47.188 64.495 63.379 63.323 1.00 30.30 A C ATOM 4745 CB HIL ARG 611 54.955 67.286 64.499 63.3823 1.00 30.30 A C ATOM 4745 CB HIL AR						FΙ	G. 4	- 97			(Conti	nued)
ATOM 4706 0E2 GLU 608 51.285 66.435 62.625 1.00 40.64 A 0 ATOM 4707 C GLU 608 50.606 63.891 60.012 1.00 32.76 A C ATOM 4708 0 GLU 608 49.889 63.527 60.947 1.00 33.47 A 0 O ATOM 4709 N ALA 609 50.643 63.270 58.836 1.00 31.32 A N ATOM 4710 CA ALA 609 49.827 62.905 58.595 1.00 30.73 A C ATOM 4711 CB ALA 609 49.827 62.905 58.595 1.00 30.73 A C ATOM 4711 CB ALA 609 49.883 61.682 57.123 1.00 28.50 A C ATOM 4713 0 ALA 609 49.883 61.682 57.123 1.00 28.50 A C ATOM 4713 1 CA ALA 609 49.583 60.274 60.139 1.00 31.03 A 0 O ATOM 4715 CA ALA 610 51.674 60.803 59.479 1.00 29.26 A N ATOM 4716 CB ALA 610 51.674 60.803 59.479 1.00 29.26 A N ATOM 4716 CB ALA 610 53.826 59.818 60.114 1.00 27.67 A C ATOM 4718 0 ALA 610 51.565 58.9818 60.174 11.00 27.67 A C ATOM 4718 0 ALA 610 51.565 58.904 62.379 1.00 28.48 A C ATOM 4719 N ARG 611 52.025 61.094 62.282 1.00 28.94 A N ATOM 4720 CA ARG 611 52.025 61.094 62.282 1.00 28.94 A N ATOM 4721 CB ARG 611 51.674 61.309 63.678 1.00 28.98 A C ATOM 4723 CD ARG 611 53.239 63.291 64.032 1.00 28.96 A C ATOM 4723 CD ARG 611 54.980 66.384 63.378 1.00 29.92 A C ATOM 4724 NE ARG 611 54.980 66.384 63.378 1.00 29.92 A C ATOM 4724 NE ARG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4728 CD ARG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CA RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CA RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CA RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CA RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CB RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CB RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CB RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CB RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CB RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CB RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CB RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CB RG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4734 CB RG 611 54.980 66.384 63.385 1.00 30.03 A C ATOM 4734 CB RG 614 63.93 38.93 38.93 39.00 30.03 A C ATOM 474	ATOM	4705	0E1	GLU	608	49.197	66.454	63.302	1.00 37.67			
ATOM 4708 0 GLU 608 49.889 63.527 60.947 1.00 33.47 A 0 ATOM 4709 N ALA 609 50.643 63.270 52.090 58.595 1.00 30.73 A C ATOM 4710 CA ALA 609 49.883 61.682 57.123 1.00 28.50 A C ATOM 4711 CB ALA 609 49.883 61.682 57.123 1.00 28.50 A C ATOM 4713 O ALA 609 49.583 60.274 60.139 1.00 31.03 A O ATOM 4713 O ALA 609 50.355 60.968 59.472 1.00 30.16 A C ATOM 4713 O ALA 609 50.355 60.968 59.472 1.00 30.16 A C ATOM 4714 N ALA 610 51.674 60.803 59.479 1.00 29.26 A N ATOM 4715 CA ALA 610 52.310 59.758 60.274 1.00 28.48 A C ATOM 4716 CB ALA 610 53.826 59.818 60.114 1.00 27.67 A C ATOM 4716 CB ALA 610 51.930 59.886 61.743 1.00 27.62 A C ATOM 4717 C ALA 610 51.930 59.886 61.743 1.00 27.62 A C ATOM 4719 N ARG 611 52.025 61.094 62.282 1.00 26.94 A N ATOM 4721 CB ARG 611 51.674 61.309 63.678 1.00 28.98 A C ATOM 4721 CB ARG 611 51.674 61.309 63.678 1.00 28.98 A C ATOM 4722 CG ARG 611 53.281 64.799 64.187 1.00 28.99 A C ATOM 4722 CD ARG 611 53.281 64.799 64.187 1.00 28.90 A C ATOM 4724 NB ARG 611 53.281 64.799 64.187 1.00 29.92 A C ATOM 4724 NB ARG 611 53.281 64.799 64.187 1.00 29.92 A C ATOM 4724 NB ARG 611 54.641 65.232 64.102 1.00 29.97 A C ATOM 4728 C ARG 611 54.946 66.384 63.378 1.00 29.97 A C ATOM 4728 C ARG 611 54.956 67.028 62.680 1.00 31.41 A N ATOM 4728 C ARG 611 54.956 67.028 62.680 1.00 31.41 A N ATOM 4730 N GLN 612 49.319 61.988 60.084 63.323 1.00 29.90 A C ATOM 4730 N GLN 612 49.319 61.988 60.084 63.323 1.00 29.97 A C ATOM 4730 N GLN 612 49.319 61.988 60.084 63.323 1.00 30.18 A N ATOM 4730 N GLN 612 49.319 61.988 60.084 61.00 31.08 A O ATOM 4730 N GLN 612 47.916 60.922 63.195 1.00 30.18 A N ATOM 4730 N GLN 612 49.319 61.988 60.084 61.00 31.08 A O A ATOM 4730 N GLN 612 47.186 64.976 63.376 63.379 63.444 1.00 35.03 A O A ATOM 4730 N GLN 612 47.186 64.976 63.376 63.379 1.00 29.90 A C ATOM 4730 N GLN 612 47.196 60.922 63.195 1.00 30.05.5 A C ATOM 4730 N GLN 612 47.196 60.922 63.195 1.00 30.05.5 A C ATOM 4730 N GLN 612 47.196 60.922 63.195 1.00 30.05.5 A C A ATOM 4744 CD PHE 613 48.835 56.679 63.597 1.00 30.42 A C A C ATOM 4												
ATOM 4710 A ALA 609												
ATOM 4710 CA ALA 609 49.827 62.000 58.595 1.00 30.73 A C ATOM 4711 CB ALA 609 49.833 61.682 57.123 1.00 28.50 A C ATOM 4712 C ALA 609 49.583 60.274 60.139 1.00 28.03 A C ATOM 4713 O ALA 609 49.583 60.274 60.139 1.00 30.16 A C ATOM 4714 N ALA 610 51.674 60.803 59.479 1.00 29.26 A N ATOM 4716 CB ALA 610 51.674 60.803 59.479 1.00 29.26 A N ATOM 4716 CB ALA 610 52.310 59.758 60.274 1.00 28.48 A C ATOM 4716 CB ALA 610 53.826 59.818 60.114 1.00 27.67 A C ATOM 4718 O ALA 610 51.566 58.904 62.379 1.00 28.48 A C ATOM 4718 O ALA 610 51.566 58.904 62.379 1.00 28.43 A O ATOM 4719 N ARG 611 52.025 61.094 62.282 1.00 26.94 A N ATOM 4720 CA ARG 611 51.674 61.309 63.678 1.00 28.43 A O ATOM 4721 CB ARG 611 51.812 62.787 64.042 1.00 28.98 A C ATOM 4721 CB ARG 611 51.812 62.787 64.042 1.00 28.96 A C ATOM 4723 CD ARG 611 53.239 63.291 64.032 1.00 29.26 A C ATOM 4724 NB ARG 611 53.239 63.291 64.032 1.00 29.26 A C ATOM 4724 NB ARG 611 54.641 65.322 64.102 1.00 29.92 A C ATOM 4726 NIH ARG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4727 NH2 ARG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4728 C ARG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4728 C ARG 611 54.980 66.384 63.378 1.00 29.97 A C ATOM 4730 N GLN 611 49.983 60.084 64.856 1.00 31.41 A N ATOM 4730 N GLN 612 49.913 60.084 64.856 1.00 31.91 A N ATOM 4730 N GLN 612 49.913 60.084 64.856 1.00 31.91 A N ATOM 4731 CA GLN 612 49.913 60.984 64.856 1.00 31.03 A O ATOM 4730 N GLN 612 49.913 60.984 64.856 1.00 31.03 A O A C ATOM 4730 N GLN 612 49.913 60.984 64.856 1.00 31.03 A O A C ATOM 4730 N GLN 612 49.913 60.984 64.856 1.00 31.91 A N ATOM 4730 N GLN 612 49.319 61.298 63.076 1.00 30.42 A C ATOM 4731 CA GLN 612 47.112 63.001 61.964 1.00 33.70 A C ATOM 4735 DEI GLN 612 47.112 63.001 61.964 1.00 33.70 A C ATOM 4734 CD GLN 612 47.128 64.475 63.379 63.441 1.00 33.70 A C ATOM 4734 CD GLN 612 47.138 64.475 63.379 63.441 1.00 33.70 A C ATOM 4744 CD PIBE 613 48.851 57.126 59.787 1.00 30.590 A N A ATOM 4744 CD PIBE 613 48.851 57.126 59.787 1.00 30.03 A C ATOM 4744 CD PIBE 613 48.855 57.126												
ATOM 4711 CB ALA 609												
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					FI	G. 4	- 98			(Continued)
ATOM	4753	0G	SER	614	52.686	57. 477		1.00 38.53	A	0
ATOM	4754	C	SER	614	49. 424	57.098		1.00 39.76	A	·C
ATOM	4755	0	SER	614	49. 283	56.276		1.00 41.47	Α	0
ATOM	4756	N	LYS	615	48. 694	58. 204		1.00 40.51	Α	N
ATOM	4757	CA	LYS	615	47.663	58.490		1.00 41.32	Α	C
ATOM	4758	CB	LYS	615	47. 047	59.870		1.00 42.73	Α	C
ATOM	4759	CG	LYS	615	47. 884			1.00 44.59	A	C
ATOM	4760	CD	LYS	615	47.064	62.330		1.00 46.18	Α	С
ATOM	4761	CE	LYS	615	47. 864	63. 511		1.00 46.73	Α	С
ATOM	4762	NZ	LYS	615	48.314	63. 301		1.00 48.03	Α	N
ATOM	4763	C	LYS	615	46.552	57. 441		1.00 40.86	Α	C
ATOM	4764	0	LYS	615	45. 794	57. 285		1.00 41.94	Α	0
ATOM	4765	N	MET	616	46.456	56.724		1.00 39.78	Α	N
ATOM	4766	CA	MET	616	45.418	55.712		1.00 37.88	Α	C
ATOM	4767	CB	MET	616	45.246	55.374	64.578	1.00 37.42	Α	C
ATOM	4768	CG	MET	616	44.673	56. 532	63.768	1.00 35.95	Α	C
ATOM	4769	SD	MET	616	44. 195	56.101	62.079	1.00 35.73	Α	S
ATOM	4770	CE	MET	616	43. 946	57.730	61.385	1.00 34.06	Α	C
ATOM	4771	C	MET	616	45.654	54.447	66.885	1.00 36.90	Α	C ·
ATOM	4772	0	MET	616	44. 908	53.473	66.772	1.00 37.22	Α	0
ATOM	4773	N	GLY	617	46.706	54.469	67.698	1.00 35.15	Α	N
ATOM	4774	CA	GLY	617	47.013	53.355	68.578	1.00 32.74	Α	C ·
MOTA	4775	C	GLY	617	47. 445	51.995	68.065	1.00 32.72	Α	C
ATOM	4776	0	GLY	617	47.806	51.143	68.872	1.00 33.71	Α	0
ATOM	4777	N	PHE	618	47. 409	51.751	66.761	1.00 32.52	Α	N
ATOM	4778	CA	PHE	618	47.841	50.447	66.262	1.00 31.36	A	С
ATOM	4779	CB	PHE	618	46.701	49.759	65.496	1.00 31.10	Α	C
ATOM	4780	CG	PHE	618	46.047	50.624	64.457	1.00 31.61	A	C
ATOM	4781	CD1	PHE	618	46.743	51.025	63. 322	1.00 31.30	A	C
ATOM	4782	CD2	PHE	618	44. 724	51.027		1.00 30.93	A	Č
ATOM	4783	CE1	PHE	618	46. 129	51.815		1.00 31.53	A	Ċ
ATOM	4784	CE2	PHE	618	44. 104	51.814		1.00 30.94	Ā	Č
ATOM	4785	CZ	PHE	618	44.808	52.209		1.00 29.86	A	Č
ATOM	4786	C	PHE	618	49.109	50.521	65.404	1.00 30.95	A	Č
ATOM	4787	.0	PHE	618	49.303	49.735	64.477	1.00 30.95	Ā	0
ATOM	4788	N	VAL	619	49. 982	51.465		1.00 30.23	A	Ň
ATOM	4789	CA	VAL	619	51.226	51.627		1.00 29.99	Ā	Ċ
ATOM	4790	CB	VAL	619	51.226	52.928		1.00 29.39	Ā	Č
ATOM	4791	CG1		619	52.632	53. 200		1.00 28.74	Ā	Č
ATOM	4792	CG2		619	50. 248	52.804		1.00 26.48	Ā	Č
ATOM	4793	C	VAL	619	52. 425	51.673	65. 931	1.00 29.66	Ä	č
ATOM	4794	•0	VAL	619	52.400	52.342	66. 962	1.00 30.05	Ä	Ö
ATOM	4795	Ň	ASP	620	53. 475	50.954		1.00 29.84	A	Ň
ATOM	4796	CA	ASP	620	54.695	50. 932	66. 347	1.00 29.07	A	Č
ATOM	4797	CB	ASP	620	55. 563	49. 748	65. 924	1.00 27.94	Ä	č
ATOM	4798	CG	ASP	620	56.789	49. 587	66. 794	1.00 27.02	Ä	č
ATOM	4799	0D1		620	57. 191	50. 580	67. 439	1.00 26.38	A	Õ
ATOM	4800	0D2		620	57. 358	48. 473	66. 818	1.00 25.22	A	0
ATOM	4801	C	ASP	620	55. 408	52. 243		1.00 20.22	A	C
111 0111	1001	U	, IUI	040	00. 400	04.410	00.000	1.00 00.00	п	v

				FIG 4-99	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	4832 4833 4834 4835 4836 4837 4838 4839 4840 4841 4842 4843 4844 4845 4846 4847 4848 4849	O ASP O ASS O ASS O ASS O D1 ASS O N CA ARG O N CA ARG O N CA ALA ALA ALA ALA ALA CB O N CA CB ILE CCD ILE O N CA CB ILE CCD ILE	621	FIG. 4 - 99 56. 009 52. 398 64. 979 1. 00 29. 95 A 55. 330 53. 196 66. 958 1. 00 33. 01 A 55. 962 54. 492 66. 746 1. 00 35. 15 A 56. 420 54. 804 69. 214 1. 00 43. 03 A 57. 648 54. 821 69. 346 1. 00 44. 79 A 55. 606 54. 280 70. 130 1. 00 45. 61 A 57. 453 54. 370 66. 441 1. 00 35. 20 A 58. 016 53. 186 66. 660 1. 00 34. 67 A 58. 016 53. 186 66. 660 1. 00 35. 70 A 60. 030 52. 027 66. 418 1. 00 37. 42 A 60. 763 51. 584 69. 804 1. 00 43. 05 A 60. 839 52. 077 71. 240 1. 00 45. 27 A 61. 516 51. 077 72. 123 1. 00 45. 73 A 62. 52. 572 64. 571 1. 00 34. 38 A	O N C C C C C C C C C C C C C C C C C C

4899

ATOM

OH

TYR

631

103/246

(Continued) FIG. 4-100 55.619 52.776 **ATOM** 4851 0 ILE 626 60.380 1.00 17.10 Α 0 58.425 55.893 51.719 1.00 17.62 N A 4852 N TRP 627 **ATOM** 50.409 C 627 58.998 55.622 1.00 17.62 A **ATOM** 4853 CA TRP 50.206 C TRP 59.190 54.118 1.00 16.80 627 Α **ATOM** 4854 CB Č TRP 627 58.096 53.441 49.427 1.00 18.70 A **ATOM** 4855 CG $_{\mathbb{C}}^{\mathbb{C}}$ 58.139 53.055 48.044 1.00 17.58 Α **ATOM** 4856 CD2 TRP 627 **ATOM** 4857 CE2 TRP 627 56.912 52.425 47.749 1.00 17.70 A 47.028 59.095 53.179 1.00 15.10 $^{\rm C}_{\rm C}$ CE3 TRP 627 A **ATOM** 4858 49.895 1.00 18.68 CD1 TRP 627 56.879 53.047 A **ATOM** 4859 N **ATOM** 4860 NE1 TRP 627 56.163 52.435 48.896 1.00 18.72 A 46.480 **ATOM** CZ2 627 56.617 51.916 1.00 16.42 A C 4861 TRP 58.801 45.769 $_{\rm C}^{\rm C}$ CZ3 TRP 627 52.673 1.00 14.48 **ATOM** 4862 Α 627 57.575 52.048 45.507 1.00 14.63 A CH2 TRP **ATOM** 4863 C 56.191 49.275 **ATOM** 4864 TRP 627 58.157 1.00 18.48 A C 56.934 49.381 0 4865 TRP 627 56.280 1.00 18.15 **ATOM** 0 A 48.193 58.829 N **ATOM** 4866 N **GLY** 628 56.579 1.00 18.70 Α 58.140 57.146 47.049 1.00 18.30 A C 4867 CA GLY 628 **ATOM** 57.163 C 58.986 45.787 1.00 18.36 ATOM 4868 C **GLY** 628 A 60.212 45.833 **ATOM** 4869 0 GLY 628 57.065 1.00 19.07 A 0 44.654 N 629 58.312 57.300 1.00 17.25 N **ATOM** 4870 TRP A 4871 CA 629 58.945 57.322 43.343 1.00 15.27 A C **ATOM** TRP Č **ATOM** 4872 CB TRP 629 58.306 56.214 42.494 1.00 10.48 A $_{\mathsf{C}}^{\mathsf{C}}$ 4873 CG TRP 629 59.131 55.698 41.357 1.00 10.84 **ATOM** A TRP 59.512 54.335 41.122 1.00 9.02 CD2 **ATOM** 4874 629 Α **ATOM** 4875 CE2 TRP 629 60.243 54.310 39.914 1.00 10.87 A C CE3 TRP 59.312 53.135 41.818 **ATOM** 4876 629 1.00 9.31 A C **ATOM** 4877 CD1 TRP 629 59.635 56, 422 40.313 1.00 10.72 A NE1 TRP 60.299 **ATOM** 4878 629 55.595 39.443 1.00 10.74 A N 629 60.779 53.126 39.379 $^{\rm C}$ 4879 CZ2 TRP 1.00 12.40 A **ATOM ATOM** 4880 CZ3 TRP 629 59.842 51.959 41.295 1.00 11.95 A C 4881 60.571 **ATOM** CH2 TRP 629 51.965 40.080 1.00 13.29 Α C 58.671 42.753 ATOM 4882 TRP 629 58.722 1.00 15.91 A 0 59.300 0 **ATOM** 4883 TRP 629 57.622 43.012 1.00 15.58 Α ATOM 4884 N SER 630 59.612 59.269 41.983 1.00 16.99 N A 4885 CA SER 630 59.453 60.603 41.383 **ATOM** 1.00 16.78 A C 58.258 4886 CB SER 630 60.644 40.421 ATOM 1.00 18.65 A 0 ATOM 4887 0G SER 630 58.531 59.987 39.198 1.00 22.38 Α 4888 SER 630 59.234 42.450 C **ATOM** C 61.656 1.00 16.69 A **ATOM** 4889 0 SER 630 60.076 61.856 43.321 1.00 17.90 0 A TYR 42.368 4890 N 631 58.093 62.335 1.00 17.21 A N ATOM 4891 CA TYR 631 57.737 63.362 43.335 C ATOM 1.00 15.51 A TYR 56.380 63.969 42.981 C **ATOM** 4892 CB 631 1.00 17.16 A Č CG 4893 TYR 631 56.161 65.353 43.545 ATOM 1.00 18.38 A 4894 CD1 TYR 631 55.947 65,550 44.909 1.00 18.79 A CCCCATOM TYR 4895 CE1 631 55.741 66.826 45.429 1.00 19.48 A ATOM CD2 TYR 631 56.168 42.714 1.00 18.85 4896 66.470 A ATOM CE2 TYR 631 55.963 43.226 1.00 19.30 **ATOM** 4897 67.751 A ATOM 4898 CZ TYR 631 55.748 67.918 44.580 1.00 19.21

69.173 SUBSTITUTE SHEET (RULE 26)

45.084

1.00 20.71

0

55.520

		FIG. 4-101	(Continued)
ATOM 4901 ATOM 4902 ATOM 4903 ATOM 4904 ATOM 4905 ATOM 4906 ATOM 4907 ATOM 4908 ATOM 4909 ATOM 4910 ATOM 4911 ATOM 4912 ATOM 4913 ATOM 4916 ATOM 4916 ATOM 4917 ATOM 4918 ATOM 4919 ATOM 4919 ATOM 4920 ATOM 4922 ATOM 4923 ATOM 4924 ATOM 4925 ATOM 4926 ATOM 4927 ATOM 4928 ATOM 4929 ATOM 4929 ATOM 4929 ATOM 4929 ATOM 4929 ATOM 4920 ATOM 4921 ATOM 4921 ATOM 4923 ATOM 4924 ATOM 4925 ATOM 4926 ATOM 4927 ATOM 4928 ATOM 4929 ATOM 4929 ATOM 4929 ATOM 4929 ATOM 4929 ATOM 4930 ATOM 4931	O TYR 634 N VAL 635 CA VAL 635 CB VAL 635 CG1 VAL 635 CG2 VAL 635 C VAL 635 C VAL 635 THR 636 CA THR 636 CB THR 636	57. 672 62. 632 44. 668 1. 00 15. 27 57. 946 63. 201 45. 731 1. 00 13. 23 57. 324 61. 350 44. 592 1. 00 14. 83 57. 266 60. 529 45. 783 1. 00 15. 04 58. 653 60. 477 46. 394 1. 00 14. 53 58. 816 60. 652 47. 596 1. 00 13. 85 59. 655 60. 246 45. 551 1. 00 15. 63 61. 030 60. 185 46. 014 1. 00 14. 69 61. 500 61. 513 46. 576 1. 00 15. 25 62. 251 61. 561 47. 555 1. 00 16. 82 61. 058 62. 598 45. 954 1. 00 13. 67 61. 418 63. 940 46. 398 1. 00 13. 29 60. 901 64. 964 45. 397 1. 00 11. 67 60. 914 66. 382 45. 904 1. 00 12. 54 62. 125 68. 388 46. 484 1. 00 13. 37 59. 723 67. 057 46. 173 1. 00 11. 38 60. 933 69. 049	(Continued) A C A 0 A N A C A C A C A C A C A C A C A C A C A C
ATOM 4933 C ATOM 4934 C ATOM 4935 O	G2 THR 636 THR 636 THR 636	59.504 59.130 49.696 1.00 20.57 61.362 58.807 51.157 1.00 20.48 61.676 61.676 51.396 1.00 19.91 61.914 61.696 52.609 1.00 19.58	A 0 A C A C A 0
ATOM 4936 N ATOM 4937 CA ATOM 4938 CI ATOM 4939 OO	A SER 637 B SER 637 G SER 637	62. 524 62. 141 50. 483 1. 00 19. 89 63. 804 62. 711 50. 862 1. 00 20. 30 64. 599 63. 086 49. 614 1. 00 19. 17 64. 823 61. 952 48. 800 1. 00 19. 07	A N A C A C A C
ATOM 4940 C ATOM 4941 0 ATOM 4942 N ATOM 4943 CA	SER 637 MET 638 A MET 638	63. 615 63. 938 51. 749 1. 00 21. 61 64. 235 64. 049 52. 812 1. 00 22. 54 62. 760 64. 855 51. 309 1. 00 21. 06 62. 490 66. 074 52. 066 1. 00 21. 87	A C A O A N A C
ATOM 4944 CE ATOM 4945 CC ATOM 4946 SL ATOM 4947 CE ATOM 4948 C	G MET 638 D MET 638 E MET 638	61. 417 66. 895 51. 354 1. 00 20. 36 61. 876 67. 465 50. 032 1. 00 21. 23 63. 069 68. 787 50. 261 1. 00 21. 33 62. 006 70. 229 50. 125 1. 00 19. 31 62. 039 65. 748 53. 494 1. 00 21. 51	A C A C A C A C A C A C A C A C

					FIG	. 4 -	102			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM	4949 4950 4951 4952 4953 4954		MET VAL VAL VAL VAL VAL	638 639 639 639 639 639	62. 511 61. 116 60. 611 59. 524 59. 201 58. 275	66. 351 64. 798 64. 372 63. 287 62. 688 63. 879	54. 472 53. 600 54. 891 54. 746 56. 112 54. 108	1.00 19.64 1.00 19.63 1.00 20.04 1.00 20.08 1.00 20.55 1.00 17.95	A A A A A	O N C C C
ATOM ATOM ATOM ATOM	4955 4956 4957 4958	C O N CA	VAL VAL LEU LEU	639 639 640 640	61. 758 61. 986 62. 489 63. 608	63. 793 64. 185 62. 864 62. 225	55. 692 56. 831 55. 088 55. 765	1.00 20.25 1.00 23.11 1.00 20.83 1.00 22.08	A A A A	C O N C
ATOM ATOM ATOM ATOM	4959 4960 4961 4962	CB CG CD1	LEU LEU LEU LEU	640 640 640 640	64. 245 63. 400 64. 143 63. 105	61. 179 59. 939 59. 041 59. 205	54. 855 54. 570 53. 611 55. 863	1.00 22.31 1.00 21.31 1.00 22.16 1.00 22.25	A A A A	C C C
ATOM ATOM ATOM ATOM	4963 4964 4965 4966	C O N CA	LEU LEU GLY GLY	640 640 641 641	64. 675 65. 416 64. 745 65. 731	63. 212 62. 922 64. 374 65. 368	56. 239 57. 182 55. 592 55. 972	1.00 23.38 1.00 22.99 1.00 23.16 1.00 23.10	A A A A	C O N C
ATOM ATOM ATOM ATOM	4967 4968 4969 4970	C O N CA	GLY GLY SER SER	641 641 642 642	65. 153 65. 782 63. 958	66. 555 67. 609 66. 393 67. 484	56. 721 56. 802 57. 278 58. 002	1.00 23.73 1.00 23.94 1.00 22.74 1.00 20.76	A A A A	C O O O O O O O O O O O O O O O O O O O
ATOM ATOM ATOM ATOM	4971 4972 4973 4974	CB OG C O	SER SER SER SER	642 642 642 642	61. 798 61. 319 63. 723 63. 656	67. 370 66. 213 67. 488 68. 519	57. 883 58. 546 59. 471 60. 140	1.00 19.77 1.00 17.97 1.00 21.73 1.00 21.40	A A A A	C 0 C 0
ATOM ATOM ATOM ATOM	4975 4976 4977 4978	N CA C O	GLY GLY GLY	643 643 643	64. 548 63. 407 63. 585	66. 327 66. 213 65. 944 66. 064	59. 967 61. 350 62. 314 63. 528	1.00 22.24 1.00 22.64 1.00 23.74 1.00 25.32	A A A	N C C O
ATOM ATOM ATOM ATOM	4979 4980 4981 4982	N CA CB OG	SER SER SER SER	644 644 644	61.067 59.850 59.898		61. 786 62. 616 61. 742 61. 247	1. 00 23. 53 1. 00 23. 38 1. 00 24. 79 1. 00 24. 45	A A A	N C C
ATOM ATOM ATOM ATOM ATOM	4983 4984 4985 4986 4987	C O N CA C	SER SER GLY GLY GLY	644 644 645 645 645	60. 565 62. 278 62. 543	64. 129 63. 961 63. 307 62. 166	63. 559 64. 536 63. 258 64. 107	1. 00 23. 18 1. 00 24. 28 1. 00 23. 27 1. 00 24. 80	A A A	C O N C
ATOM ATOM ATOM ATOM	4988 4989 4990 4991	O N CA CB	GLY VAL VAL VAL	645 646 646 646	61. 379 60. 446 59. 289	61. 175 60. 248 61. 357 60. 474 61. 207	64. 114 64. 920 63. 207 63. 121 62. 473	1. 00 24. 80 1. 00 27. 93 1. 00 23. 98 1. 00 22. 32 1. 00 24. 36	A A A A	C O N C
ATOM ATOM ATOM ATOM	4992 4993 4994 4995	CG1 CG2 C	VAL	646 646 646 646	56. 945 57. 636 59. 552	60. 230 62. 351 59. 202 58. 128	62. 215 63. 381 62. 327 62. 690	1. 00 24. 30 1. 00 22. 37 1. 00 24. 11 1. 00 21. 28 1. 00 21. 25	A A A A	C C C O
ATOM ATOM	4996 4997	N	PHE PHE	647 647	60. 303	59. 326 58. 182	61. 239 60. 380	1. 00 21. 20 1. 00 21. 00 1. 00 18. 33	A A	N C

		FIG. 4-103	(Continued)
ATOM 4998 CB PHE ATOM 4999 CG PHE ATOM 5000 CD1 PHE ATOM 5001 CD2 PHE ATOM 5003 CE2 PHE ATOM 5003 CE2 PHE ATOM 5006 C PHE ATOM 5006 C PHE ATOM 5007 N LYS ATOM 5008 CA LYS ATOM 5010 CG LYS ATOM 5010 CG LYS ATOM 5011 CD LYS ATOM 5012 CE LYS ATOM 5012 CE LYS ATOM 5013 NZ LYS ATOM 5014 C LYS ATOM 5015 C LYS ATOM 5016 N CYS ATOM 5017 CA CYS ATOM 5018 C CYS ATOM 5018 C CYS ATOM 5019 CYS ATOM 5019 CYS ATOM 5020 CB CYS ATOM 5020 CB CYS ATOM 5021 CC CYS ATOM 5021 CC CYS ATOM 5020 CB CYS ATOM 5020 CB CYS ATOM 5021 CC CYS ATOM 5021 CC CYS ATOM 5022 N GLY ATOM 5023 CA GLY ATOM 5024 C GLY ATOM 5025 C GLY ATOM 5026 N ILE ATOM 5027 CA ILE ATOM 5028 CB ILE ATOM 5029 CG2 ILE ATOM 5030 CG1 ILE ATOM 5031 CD1 ILE ATOM 5032 C ILE ATOM 5033 C ILE ATOM 5034 N ALA ATOM 5035 CA ALA ATOM 5036 CB ALA ATOM 5037 C ALA ATOM 5038 C ALA ATOM 5039 N VAL ATOM 5039 N VAL ATOM 5041 CB VAL ATOM 5042 CG1 VAL	647 647 647 647 647 647 647 648 648 648 648 648 648	60. 497 58. 615 58. 924 1. 00 15. 79 A 59. 142 59. 131 58. 551 1. 00 16. 11 A 58. 138 58. 258 58. 152 1. 00 15. 39 A 58. 841 60. 479 58. 680 1. 00 14. 43 A 56. 855 58. 722 57. 894 1. 00 13. 82 A 57. 562 60. 943 58. 423 1. 00 15. 28 A 56. 568 60. 061 58. 031 1. 00 13. 75 A 61. 944 57. 555 60. 663 1. 00 18. 46 A 62. 943 58. 250 60. 825 1. 00 20. 84 A 61. 958 56. 232 60. 722 1. 00 17. 11 A 63. 165 55. 480 60. 996 1. 00 19. 06 A 62. 789 54. 105 61. 545 1. 00 17. 86 A 63. 961 53. 242 61. 955 1. 00 17. 94 A 63. 484 51. 869 62. 405 1. 00 19. 57 A 64. 594 51. 083 63. 095 1. 00 19. 22 A 65. 757 50. 894 62. 204 1. 00 20. 59 A 64. 025 55. 314 59. 747 1. 00 21. 47 A 65. 251 55. 379 59. 815 1. 00 23. 13 A 63. 376 55. 094 58. 610 1. 00 22. 38 A 64. 077 54. 898 57. 353 1. 00 24. 23 A 63. 156 55. 237 56. 181 1. 00 24. 23 A 63. 156 55. 237 56. 181 1. 00 24. 09 A 61. 939 55. 319 56. 342 1. 00 23. 94 A 64. 527 53. 447 57. 237 1. 00 27. 68 A 63. 130 52. 287 57. 313 1. 00 24. 23 A 63. 376 55. 094 58. 610 0. 22. 38 A 64. 077 54. 898 57. 353 1. 00 24. 23 A 63. 130 52. 287 57. 313 1. 00 21. 50 A 62. 961 55. 757 53. 834 1. 00 21. 04 A 63. 649 55. 384 52. 535 1. 00 21. 13 A 64. 874 55. 333 52. 474 1. 00 21. 62 A 62. 857 55. 124 51. 499 1. 00 19. 35 A 63. 388 54. 753 50. 195 1. 00 19. 18 A 62. 896 53. 352 49. 758 1. 00 19. 35 A 63. 768 56. 015 48. 949 1. 00 19. 53 A 64. 323 58. 574 47. 594 1. 00 19. 53 A 64. 323 58. 574 47. 594 1. 00 19. 53 A 64. 323 58. 574 47. 594 1. 00 19. 53 A 64. 323 58. 574 47. 594 1. 00 19. 53 A 64. 323 58. 574 47. 594 1. 00 19. 53 A 64. 323 58. 574 47. 594 1. 00 19. 50 A 65. 288 56. 662 45. 996 1. 00 14. 98 A 662. 582 54. 754 48. 393 1. 00 14. 98 A 663. 692 55. 963 48. 393 1. 00 14. 84 A 62. 586 54. 291 41. 932 1. 00 14. 95 A	Continued) C C C C C C C C C C C C C C C C C C C
ATOM 5043 CG2 VAL ATOM 5044 C VAL ATOM 5045 O VAL ATOM 5046 N ALA	653 653 653 654	62. 806 53. 607 44. 352 1. 00 18. 10 A 63. 292 57. 063 42. 694 1. 00 13. 22 A 62. 224 57. 669 42. 620 1. 00 11. 12 A 64. 331 57. 317 41. 901 1. 00 12. 68 A	C C O N

		(Continued)				
ATOM	5047		LA 654		A	С
ATOM	5048		LA 654	63. 513 57. 790 39. 650 1. 00 7. 27	Α	C
ATOM	5049		LA 654		Α	C
ATOM	5050		LA 654	62.687 60.103 40.787 1.00 13.18	A	.0
ATOM	5051		RO 655	64. 208 60. 179 42. 420 1. 00 10. 68	Ą	Ŋ
ATOM ATOM	5052		RO 655	65. 319 59. 696 43. 262 1. 00 8. 01	A	C
ATOM	5053 5054		RO 655	63.643 61.408 42.971 1.00 10.40	A	C
ATOM	5055		RO 655 RO 655	64. 092 61. 344 44. 422 1. 00 8. 50	A	C
ATOM	5056		RO 655	65.476 60.822 44.277 1.00 6.23 64.090 62.714 42.327 1.00 12.92	A	C
ATOM	5057		RO 655		A	C
ATOM	5058		AL 656	65. 166 62. 793 41. 717 1. 00 13. 38 63. 245 63. 735 42. 454 1. 00 12. 39	A	0
ATOM	5059		AL 656	63. 612 65. 065 41. 999 1. 00 12. 85	A	N
ATOM	5060		AL 656	62. 373 65. 946 41. 769 1. 00 11. 42	A A	C
ATOM	5061	CG1 V		62.781 67.416 41.645 1.00 10.52	A	C C
ATOM	5062	CG2 V		61.661 65.500 40.510 1.00 10.18	A	Č
ATOM	5063		AL 656	64. 382 65. 560 43. 236 1. 00 13. 79	A	C
ATOM	5064	0 V		64. 038 65. 188 44. 355 1. 00 14. 63	Λ A	Ö
ATOM	5065		ER 657	65. 419 66. 372 43. 066 1. 00 14. 27	A	N
ATOM	5066	CA SI		66. 174 66. 831 44. 238 1. 00 14. 99	Ä	Č
ATOM	5067	CB SF	ER 657	67. 589 66. 231 44. 231 1. 00 15. 67	Ä	. Č
ATOM	5068	OG SE		68. 385 66. 819 43. 213 1. 00 15. 19	Ä	Ö
ATOM	5069	C SE		66. 286 68. 343 44. 320 1. 00 14. 77	Ä	Č
ATOM	5070	0 SE		66. 387 68. 912 45. 406 1. 00 14. 39	A	Ö
ATOM	5071	N AF		66. 269 68. 978 43. 158 1. 00 15. 05	Α	N
ATOM	5072	CA AF		66.388 70.423 43.038 1.00 16.33	Α	С
ATOM	5073	CB AF		67. 845 70. 787 42. 747 1. 00 20. 44	Α	C
ATOM	5074	CG AF		68. 142 72. 274 42. 582 1. 00 24. 34	Α	C
ATOM	5075	CD AF		69. 543 72. 450 42. 025 1. 00 25. 38	Α	C
ATOM ATOM	5076	NE AR	_	69. 905 73. 838 41. 757 1. 00 25. 70	Α	N
ATOM	5077 5078	CZ AR NH1 AR		70. 353 74. 683 42. 676 1. 00 28. 34	Α	·C
ATOM	5079	NH2 AR		70. 491 74. 288 43. 935 1. 00 28. 23	A	N
ATOM	5080	C AR		70. 690 75. 916 42. 329 1. 00 29. 55	A	N
ATOM	5081		_	65.515 70.775 41.850 1.00 15.87 65.752 70.288 40.735 1.00 16.75	A	C
ATOM	5082	N TR			A	0
ATOM	5083	CA TR		64.514 71.616 42.073 1.00 13.52 63.603 71.967 40.999 1.00 13.69	A	N
ATOM	5084	CB TR		62. 465 72. 823 41. 550 1. 00 13. 63	A	C
ATOM	5085	CG TR		61.504 71.963 42.341 1.00 17.48	A A	C
ATOM	5086	CD2 TR		60.690 70.898 41.829 1.00 16.63	A	C
ATOM	5087	CE2 TR		60. 027 70. 313 42. 927 1. 00 18. 08	A	C
ATOM	5088	CE3 TR		60. 460 70. 382 40. 547 1. 00 16. 21	A	C
ATOM	5089	CD1 TR		61.300 71.980 43.692 1.00 17.21	A	C
ATOM	5090	NE1 TR	P 659	60.418 70.993 44.050 1.00 17.37	Ä	N
ATOM	5091	CZ2 TR	P 659	59. 145 69. 233 42. 785 1. 00 21. 55	Ä	Č
ATOM	5092	CZ3 TR		59. 584 69. 311 40. 403 1. 00 18. 00	Ä	č
ATOM	5093	CH2 TR	,	58.937 68.746 41.516 1.00 20.15	A	Č
ATOM	5094	C TR		64. 219 72. 580 39. 748 1. 00 13. 15	Α	Č
ATOM	5095	O. TR	P 659	63. 643 72. 503 38. 670 1. 00 11. 17	Α	0

				FΙ	G. 4	- 105	5		(Continu	ıed)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5102 OI 5103 C 5104 O 5105 N 5106 C 5107 CI 5108 CC 5109 CI 5110 CE 5111 CI 5112 CE 5113 CZ 5114 OH 5115 C 5116 O 5117 N 5118 CA 5119 CB 5120 CG 5121 CD 5122 CE 5123 CD 5124 CE 5125 CZ 5126 OH 5127 C 5128 O 5127 C 5128 O 5127 C 5128 CZ 5128 CZ 5129 N 5130 CA 5131 CB 5132 CG 5131 CB 5132 CG 5133 OD	G GLU TYR G TYR	660 660 660 660 660 661 661 661 661 661	F I 65. 400 66. 042 67. 147 66. 548 67. 535 68. 310 67. 527 66. 577 67. 001 66. 539 67. 642 68. 878 69. 743 70. 338 71. 183 72. 322 65. 842 66. 077 64. 602 63. 445 62. 305 61. 395 60. 010 59. 184 61. 930 61. 122 59. 756 58. 983 62. 964 63. 320 61. 686 60. 998 59. 686 59. 756 58. 983 60. 945 60. 945 60. 945 58. 715 58. 234	73. 163 73. 725 74. 704 8 76. 001 76. 901 77. 600 77. 635 72. 922 71. 383 70. 269 69. 154 69. 504 70. 531 70. 805 68. 765 69. 027 70. 041 70. 252 69. 637	39. 871 38. 697 39. 108 39. 626 40. 313 39. 617 41. 561 37. 777 36. 659 38. 233 37. 399 38. 230	1.00 14.12 1.00 15.96 1.00 16.83 1.00 19.65 1.00 22.71 1.00 25.18 1.00 23.59 1.00 15.29 1.00 16.67 1.00 14.54 1.00 14.57	A A A A A A A A A A A A A A A A A A A	(Continue N C C C C C C C C C C C C C C C C C C	
ATOM ATOM	5143 N 5144 CA	VAL VAL	665 665	57. 987 56. 540	73. 247 73. 101	31.658 31.733	1.00 13.43 1.00 14.34	A A	N C	

					FI	G. A.	- 107			(Continued)
						O. 4	107			
ATOM	5194		TYR		53. 321	70. 985	38. 537	1.00 22.18	Α	С
ATOM	5195		TYR	670	51.726		39.850	1.00 19.78	A	С
ATOM	5196		2 TYR	670	51.139		39.079	1.00 19.87	A	С
ATOM	5197	CZ	TYR	670	51.944		38. 422	1.00 22.17	A	C
ATOM	5198	OH	TYR	670	51.388		37.623	1.00 23.11	A	0
ATOM	5199	C	TYR	670	54.769	76. 317	40.757	1.00 23.32	A	С
ATOM	5200	0	TYR	670	54.442	76.937	41.763	1.00 24.86	A	0
ATOM	5201	N	MET	671	55.983	76.404	40. 228	1.00 24.66	A	N
ATOM	5202	CA	MET	671	57.029	77. 207	40.851	1.00 23.96	A	·C
ATOM	5203	CB	MET	671	58. 327	76.400	40.905	1.00 24.00	A	C .
ATOM	5204	CG	MET	671	58. 288	75. 215	41.852	1.00 23.55	A	С
ATOM	5205	SD	MET	671	58. 383	75. 732	43. 565	1.00 24.97	A	S
ATOM	5206	CE	MET	671	60.159	75. 998	43. 721	1.00 21.94	A	C
ATOM	5207	C	MET	671	57. 330	78. 547	40. 203	1.00 24.00	A	C
ATOM	5208	0	MET	671	58. 101	79. 331	40. 756	1.00 25.98	A	0
ATOM	5209	N	GLY	672	56. 741	78. 822	39. 045	1.00 22.07	A	N
ATOM ATOM	5210	CA	GLY	672	57.044	80.076	38. 379	1.00 22.40	Ą	C
ATOM	5211 5212	C	GLY	672	58. 472	80.028	37. 857	1.00 22.69	A	C
ATOM	5212	O N	GLY LEU	672	59.005	78. 947	37.641	1.00 23.27	A	0
ATOM	5214	CA	LEU	673 673	59.108	81. 180	37.667	1.00 22.65	A	N
ATOM	5215	CB	LEU	673	60. 477 60. 626	81.209	37. 151	1.00 20.90	A	C
ATOM	5216	CG	LEU	673	59.639	82. 356	36. 164	1.00 19.50	A	C
ATOM	5217		LEU	673	59.779	82. 282 83. 513	35. 010 34. 147	1.00 19.96	A	C
ATOM	5218		LEU	673	59.892	81. 027	34. 203	1.00 20.87 1.00 21.63	A	C .
ATOM	5219	C	LEU	673	61.528	81. 344	38. 248	1.00 21.03	A A	C C
ATOM	5220	Ŏ	LEU	673	61.313	82. 028	39. 239	1.00 21.87	A	0
ATOM	5221	N	PRO	674	00 000	80. 700	38. 072	1.00 21.00	A	N
ATOM	5222	CD	PRO	674	63.050	79. 803	36. 968	1.00 21.16	A	Č
ATOM	5223	CA	PRO	674	63.780	80. 747	39.050	1.00 23.23	A	Č
ATOM	5224	CB	PR ₀	674	64.618	79. 510	38. 709	1.00 21.90	A	č
ATOM	5225	CG	PRO	674	63.803	78. 755	37.695	1.00 22.34	Ä	č
ATOM	5226	C	PRO	674	64.617	82.023	38.943	1.00 24.90	Ä	č ·
ATOM	5227	0	PR0	674	65.841	81.977	39.028	1.00 26.10	Ä	Ö
ATOM	5228	N	THR	675	63.966	83. 158	38.743	1.00 25.88	A	N
ATOM	5229	CA	THR	675	64.695	84.411	38.640	1.00 27.60	Α	C
ATOM	5230	CB	THR	675	64. 208	85. 237	37.447	1.00 27.12	A	C
ATOM	5231	0G1	THR	675	62.811	85. 524	37.599	1.00 29.30	Α	0
ATOM	5232		THR	675	64.431	84. 471	36. 156	1.00 25.59	Α	С
ATOM	5233	C	THR	675	64.496	85. 211	39.918	1.00 28.74	Α	C
ATOM	5234	0	THR	675	63. 543	84. 982	40.660	1.00 29.47	Α	0
ATOM	5235	N	PRO	676 ⁻	65.404	86. 156	40.200	1.00 29.41	Α	N
ATOM	5236	CD	PRO	676	66. 625	86.508	39.457	1.00 28.96	A	C
ATOM	5237	CA	PRO	676	65. 284	86.969	41.411	1.00 29.70	A	C
ATOM	5238	CB	PRO	676	66.465	87. 929	41. 299	1.00 28.87	A	C
ATOM	5239	CG	PRO	676	67.467	87. 142	40. 533	1.00 28.27	A	Č ·
ATOM	5240	C	PRO	676	63. 948	87. 707	41.484	1.00 30.03	A	C
ATOM	5241	0 M	PRO	676	63. 359	87. 829	42.558	1.00 29.93	A	0
ATOM	5242	N	GLU	677	63.463	88. 190	40. 343	1.00 30.62	A	N

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				FIG. 4-108	(Continued)
		.4		110. 4 100	
ATOM	5243	CA GLU	677	62. 203 88. 923 40. 348 1. 00 30. 92 A	С
ATOM	5244	CB GLU	677	62. 192 90. 013 39. 264 1. 00 32. 38 A	č
ATOM	5245	CG GLU	677	62. 103 89. 536 37. 821 1. 00 34. 78 A	č
ATOM	5246	CD GLU	677	63. 380 88. 877 37. 331 1. 00 37. 04 A	č
ATOM	5247	OE1 GLU	677	64. 480 89. 356 37. 697 1. 00 35. 11 A	0
ATOM	5248	0E2 GLU	677	63. 276 87. 891 36. 566 1. 00 37. 80 A	0
ATOM	5249	C GLU	677	60. 952 88. 065 40. 231 1. 00 30. 10 A	С
ATOM	5250	0 GLU	677	59.893 88.564 39.849 1.00 31.67 A	0
ATOM	5251	N ASP	678	61. 067 86. 777 40. 546 1. 00 28. 40 A	. N
ATOM	5252	CA ASP	678	59. 906 85. 897 40. 523 1. 00 26. 09 A	C
ATOM	5253	CB ASP	678	59. 833 85. 048 39. 253 1. 00 25. 88 A	C
ATOM	5254	CG ASP	678	58. 472 84. 359 39. 097 1. 00 28. 22 A	C
MOTA	5255	OD1 ASP	678	57. 885 83. 980 40. 128 1. 00 28. 64 A	0
ATOM	5256	OD2 ASP	678	57. 980 84. 189 37. 956 1. 00 28. 80 A	0
MOTA	5257 5258	C ASP O ASP	678 678	59. 920 84. 982 41. 737 1. 00 25. 86 A 59. 481 85. 382 42. 810 1. 00 28. 55 A	C
ATOM ATOM	5259	O ASP N ASN	679	59. 481 85. 382 42. 810 1. 00 28. 55 A 60. 442 83. 768 41. 591 1. 00 23. 97 A	O N
ATOM	5260	CA ASN	679	60.442 83.708 41.331 1.00 23.37 A	C .
ATOM	5261	CB ASN	679	59. 326 81. 818 42. 496 1. 00 19. 41 A	Č
ATOM	5262	CG ASN	679	58. 894 81. 146 43. 778 1. 00 19. 58 A	č
ATOM	5263	OD1 ASN	679	58. 491 79. 981 43. 775 1. 00 20. 44 A	ŏ
ATOM	5264	ND2 ASN	679	58. 957 81. 879 44. 882 1. 00 18. 70 A	Ň .
ATOM	5265	C ASN	679	61.760 82.099 42.957 1.00 21.79 A	Ċ
ATOM	5266	0 ASN	679.	61.770 81.055 43.601 1.00 21.89 A	0
ATOM	5267	N LEU	680	62. 873 82. 636 42. 472 1. 00 24. 38 A	N .
ATOM	5268	CA LEU	680	64.164 81.967 42.665 1.00 26.33 A	C
ATOM	5269	CB LEU	680	65. 316 82. 842 42. 157 1. 00 26. 74 A	C
ATOM	5270	CG LEU	680	66. 726 82. 275 42. 385 1. 00 28. 22 A	C
ATOM	5271	CD1 LEU	680	66.844 80.903 41.747 1.00 30.03 A	C
ATOM	5272 5273	CD2 LEU	680	67. 772 83. 211 41. 801 1. 00 29. 33 A	C
ATOM ATOM	5274	C LEU O LEU	680 680	64. 449 81. 556 44. 109 1. 00 27. 18 A 64. 977 80. 471 44. 347 1. 00 28. 31 A	C
ATOM	5275	N ASP	681	64. 977 80. 471 44. 347 1. 00 28. 31 A 64. 111 82. 411 45. 072 1. 00 27. 79 A	0 N
ATOM	5276	CA ASP	681	64. 360 82. 091 46. 475 1. 00 28. 03 A	C
ATOM	5277	CB ASP	681	63.836 83.196 47.394 1.00 30.36 A	Č
ATOM	5278	CG ASP	681	64.774 84.386 47.473 1.00 34.23 A	č
ATOM	5279	OD1 ASP	681	65. 908 84. 289 46. 952 1. 00 35. 59 A	ŏ
ATOM	5280	OD2 ASP	681	64. 380 85. 417 48. 067 1. 00 36. 71 A	Ŏ
ATOM	5281	C ASP	681	63. 773 80. 753 46. 920 1. 00 27. 55 A	Č
ATOM	5282	O ASP	681	64. 428 80. 005 47. 647 1. 00 28. 05 A	0
ATOM	5283	N HIS	682	62. 551 80. 438 46. 502 1. 00 25. 37 A	N
ATOM	5284	CA HIS	682	61. 981 79. 164 46. 913 1. 00 25. 07 A	C
ATOM	5285	CB HIS	682	60. 456 79. 161 46. 801 1. 00 25. 14 A	C
ATOM	5286	CG HIS	682	59. 832 77. 914 47. 349 1. 00 27. 18 A	C
ATOM	5287	CD2 HIS	682	59. 091 76. 948 46. 754 1. 00 27. 87 A	C
ATOM	5288 5280	ND1 HIS	682	60. 021 77. 503 48. 650 1. 00 26. 29 A	N
ATOM ATOM	5289 5290	CE1 HIS NE2 HIS	682 682	59. 428 76. 336 48. 832 1. 00 26. 61 A	C
ATOM	5290 5291	C HIS	682	58. 857 75. 977 47. 697 1. 00 25. 03 A 62. 559 77. 983 46. 130 1. 00 24. 30 A	N C
VIOM	ULJI	0 1110	004	62.559 77.983 46.130 1.00 24.30 A	C

ATOM 5292 0 HIS 682 62.463 76.837 46.572 1.00 23.47 A 0 ATOM 5293 K TYR 683 63.144 78.258 44.966 1.00 23.49 A N ATOM 5294 CA TYR 683 63.768 77.208 44.157 1.00 22.64 A C ATOM 5295 CB TYR 683 63.291 77.594 41.655 1.00 19.28 A C ATOM 5295 CB TYR 683 63.291 77.594 41.655 1.00 19.28 A C ATOM 5296 CD TYR 683 63.291 77.594 41.655 1.00 19.28 A C ATOM 5299 CD TYR 683 63.291 77.594 41.655 1.00 19.28 A C ATOM 5299 CD TYR 683 62.291 77.594 41.655 1.00 19.28 A C ATOM 5299 CD TYR 683 62.361 78.589 41.347 1.00 20.47 A C ATOM 5299 CD TYR 683 62.361 78.589 41.347 1.00 20.47 A C ATOM 5300 CE2 TYR 683 51.495 78.453 40.276 1.00 20.47 A C ATOM 5301 CZ TYR 683 61.495 78.453 40.276 1.00 20.47 A C ATOM 5302 OH TYR 683 65.89 77.553 45.200 1.00 19.09 A C ATOM 5303 C TYR 683 65.189 75.533 45.125 1.00 22.32 A C ATOM 5304 0 TYR 683 65.189 75.533 45.125 1.00 22.32 A C ATOM 5305 N ARG 684 65.799 77.685 45.355 1.00 22.265 A O ATOM 5306 CA ARG 684 67.025 77.392 46.076 1.00 22.97 A C ATOM 5307 CB ARG 684 69.233 78.020 44.004 1.00 22.31 A C ATOM 5308 C ARG 684 69.233 78.223 42.562 1.00 22.457 A C ATOM 5309 C DARG 684 69.237 78.28 78.223 42.562 1.00 22.47 A N ATOM 5310 NE ARG 684 69.338 78.223 42.562 1.00 23.11 A C ATOM 5311 C ARG 684 69.338 78.223 42.562 1.00 23.11 A C ATOM 5314 C ARG 684 69.338 78.223 42.562 1.00 23.11 A C ATOM 5310 NE ARG 684 69.338 78.223 42.562 1.00 23.14 A N ATOM 5311 C ARG 684 69.338 78.223 42.562 1.00 24.57 A C ATOM 5310 NE ARG 684 69.338 78.223 42.562 1.00 23.14 A N ATOM 5311 C ARG 684 69.338 78.223 42.562 1.00 23.17 A N ATOM 5310 NE ARG 684 69.338 78.223 42.562 1.00 23.17 A N ATOM 5310 NE ARG 684 69.346 79.388 40.648 1.10 20.90 A C ATOM 5320 CD ARG 684 69.446 69.47 71.71 76.368 48.111 1.00 24.16 A N ATOM 5310 NE ARG 684 69.47 77.17 78.38 40.07 1.00 29.09 A C ATOM 5320 CD ARG 684 67.71 78.38 40.08 1.00 24.14 A C ATOM 5331 C A BR 686 64.69 77.71 78.38 40.07 1.00 23.72 A C ATOM 5320 CD ARG 684 69.23 77.831 60.74 1.00 19.20 A C ATOM 5331 C A BR 686 64.69 77.71 78.38 40.00 10.00 19.20 A C ATOM 5331 C A BR 686 64.69 77.71 78.						FΙ	G. 4	109			(Continue	d)
ATOM 5294 CA TYR 683 63.768 77.208 44.157 1.00 22.64 A C ATOM 5295 CB TYR 683 64.249 77.758 42.812 1.00 20.68 A C ATOM 5296 CG TYR 683 63.291 77.7594 41.655 1.00 19.28 A C ATOM 5297 CDI TYR 683 63.325 76.461 40.857 1.00 16.29 A C ATOM 5298 CEI TYR 683 62.361 78.589 41.347 1.00 20.47 A C ATOM 5299 CD2 TYR 683 62.361 78.589 41.347 1.00 20.47 A C ATOM 5300 CZ TYR 683 61.495 78.453 40.276 1.00 20.17 A C ATOM 5301 CZ TYR 683 61.554 77.314 39.500 1.00 19.09 A C ATOM 5301 CZ TYR 683 61.554 77.314 39.500 1.00 19.09 A C ATOM 5303 C TYR 683 66.554 77.314 39.500 1.00 19.09 A C ATOM 5303 C TYR 683 66.554 77.316 38.441 1.00 21.54 A O ATOM 5303 C TYR 683 66.554 77.392 46.076 1.00 22.32 A C ATOM 5305 C TYR 683 66.95 77.176 38.441 1.00 21.54 A O ATOM 5306 CA ARG 684 67.025 77.392 46.076 1.00 22.39 A C ATOM 5307 CB ARG 684 67.025 77.392 46.076 1.00 22.44 A N ATOM 5308 C ARG 684 67.928 78.624 46.071 1.00 22.89 A C ATOM 5309 CD ARG 684 67.928 78.624 46.071 1.00 22.45 A O ATOM 5309 CD ARG 684 69.328 78.223 42.562 1.00 25.47 A N ATOM 5311 CZ ARG 684 69.328 78.223 42.562 1.00 25.47 A N ATOM 5312 NILL ARG 684 69.328 78.223 42.562 1.00 25.47 A N ATOM 5311 CZ ARG 684 66.938 79.294 42.703 1.00 29.09 A N ATOM 5312 NILL ARG 684 69.328 78.223 42.562 1.00 25.47 A N ATOM 5312 NILL ARG 684 69.328 78.223 42.562 1.00 25.47 A N ATOM 5312 NILL ARG 684 66.807 76.922 47.501 1.00 22.90 A C ATOM 5310 NE ARG 684 66.807 76.922 47.501 1.00 22.90 A N ATOM 5310 NE ARG 684 66.807 76.922 47.501 1.00 22.90 A N ATOM 5310 NE ARG 684 66.807 76.922 47.501 1.00 22.90 A N ATOM 5310 NE ARG 684 66.807 76.922 47.501 1.00 22.90 A N ATOM 5310 NE ARG 684 66.807 76.929 50.649 1.00 23.72 A C ATOM 5310 NE ARG 684 66.807 76.929 50.649 1.00 23.72 A C ATOM 5310 NE ARG 684 66.807 76.929 50.649 1.00 23.72 A C ATOM 5310 NE ARG 684 66.807 76.929 50.649 1.00 23.72 A C ATOM 5310 NE ARG 684 66.807 76.929 50.649 1.00 23.72 A C ATOM 5310 NE ARG 684 66.807 76.929 50.00 40.00 1.00 12.10 A C ATOM 5320 DDI ANN 685 65.608 77.117 78.368 48.111 1.00 24.16 A O ATOM 5331 CA AND 685 66.450 77.5									1.00 23.47	Α	0	
ATOM 5295 CB TYR 683 63.291 77.758 42.812 1.00 20.88 A C ATOM 5296 CC TYR 683 63.291 77.594 41.655 1.00 19.28 A C ATOM 5297 CDI TYR 683 63.291 77.594 41.655 1.00 19.28 A C ATOM 5299 CD2 TYR 683 62.361 78.589 41.347 1.00 20.47 A C ATOM 5299 CD2 TYR 683 62.361 78.589 41.347 1.00 20.47 A C ATOM 5290 CD2 TYR 683 61.495 78.453 40.276 1.00 19.98 A C ATOM 5290 CD2 TYR 683 61.495 78.453 40.276 1.00 20.17 A C ATOM 5301 CZ TYR 683 61.495 78.453 40.276 1.00 20.17 A C ATOM 5301 CZ TYR 683 61.554 77.314 39.500 1.00 19.09 A C ATOM 5302 OH TYR 683 60.695 77.176 38.441 1.00 21.54 A O ATOM 5303 C TYR 683 65.189 75.533 45.125 1.00 22.65 A O ATOM 5305 N ARG 684 67.025 77.392 46.076 1.00 22.25 A C ATOM 5306 CA ARG 684 67.025 77.392 46.076 1.00 22.267 A C ATOM 5307 C BARG 684 67.025 77.392 46.076 1.00 22.97 A C ATOM 5308 C ARG 684 67.928 78.624 46.071 1.00 22.89 A C ATOM 5308 C ARG 684 69.338 78.020 44.004 1.00 23.11 A C ATOM 5301 C ARG 684 69.348 79.299 41.974 1.00 27.89 A C ATOM 5310 NE ARG 684 69.344 79.299 41.974 1.00 27.89 A C ATOM 5311 CZ ARG 684 69.344 79.299 41.974 1.00 27.89 A C ATOM 5311 CZ ARG 684 69.344 79.299 41.974 1.00 27.89 A C ATOM 5312 NII ARG 684 69.346 79.388 40.648 1.00 27.04 A N ATOM 5312 NII ARG 684 69.346 79.388 40.648 1.00 27.04 A N ATOM 5316 N ASN 685 65.608 77.17 6.384 41.11 1.00 24.16 A N ATOM 5316 N ASN 685 65.608 77.17 46.389 41.11 1.00 24.16 A N ATOM 5316 N ASN 685 65.608 77.17 46.389 41.11 1.00 24.16 A N A ATOM 5319 C ASN 685 64.455 77.547 51.610 1.00 32.842 A C ATOM 5321 ND ASN 685 66.456 77.79 15 22.160 1.00 37.49 A N A C ATOM 5321 ND ASN 685 66.465 77.79 15 22.666 1.00 38.25 A O A C ATOM 5321 ND ASN 685 66.465 77.79 15 22.160 1.00 37.49 A N A C ATOM 5321 ND ASN 685 66.465 77.79 15 22.666 1.00 38.25 A O A ATOM 5322 C ASN 685 64.456 77.11 76.368 48.11 1.00 24.16 A N A N A ATOM 5322 C ASN 685 64.456 77.11 76.368 48.11 1.00 23.37 2 A C C ATOM 5323 ND ASN 685 66.406 77.79 15 22.666 1.00 13.74 A C A ATOM 5322 C ASN 685 64.456 77.79 15 22.606 1.00 17.15 A C ATOM 5323 C A SR 686 64.101 74.852 48.417 1.00										Α		
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ATOM 5323 O ASN 685 64.356 74.929 50.649 1.00 23.86 A O ATOM 5324 N SER 686 64.101 74.852 48.417 1.00 21.55 A N ATOM 5325 CA SER 686 63.336 73.613 48.457 1.00 19.71 A C ATOM 5326 CB SER 686 61.976 73.811 47.774 1.00 19.20 A C ATOM 5327 OG SER 686 62.114 74.112 46.397 1.00 15.00 A O ATOM 5328 C SER 686 64.060 72.421 47.823 1.00 20.13 A C ATOM 5329 O SER 686 63.447 71.611 47.128 1.00 21.27 A O ATOM 5330 N THR 687 65.362 72.307 48.060 1.00 19.02 A N ATOM 5331 CA THR 687 66.122 71.189 47.509 1.00 17.15 A C ATOM 5332 CB THR 687 67.441 71.665 46.906 1.00 19.02 A N ATOM 5333 OG1 THR 687 68.362 71.959 47.960 1.00 17.42 A O ATOM 5334 CG2 THR 687 67.441 71.665 46.906 1.00 16.10 A C ATOM 5335 C THR 687 66.433 70.153 48.585 1.00 15.79 A C ATOM 5336 O THR 687 66.433 70.153 48.585 1.00 15.79 A C ATOM 5337 N VAL 688 66.496 70.466 49.763 1.00 15.82 A O ATOM 5337 N VAL 688 66.627 68.908 48.182 1.00 18.43 A N ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5338 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C		5322										
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ATOM 5328 C SER 686 64.060 72.421 47.823 1.00 20.13 A C ATOM 5329 0 SER 686 63.447 71.611 47.128 1.00 21.27 A 0 ATOM 5330 N THR 687 65.362 72.307 48.060 1.00 19.02 A N ATOM 5331 CA THR 687 66.122 71.189 47.509 1.00 17.15 A C ATOM 5332 CB THR 687 67.441 71.665 46.906 1.00 16.10 A C ATOM 5333 0G1 THR 687 68.362 71.959 47.960 1.00 17.42 A 0 ATOM 5334 CG2 THR 687 67.214 72.920 46.058 1.00 14.71 A C ATOM 5335 C THR 687 66.433 70.153 48.585 1.00 14.71 A C ATOM 5336 0 THR 687 66.496 70.466 49.763 1.00 15.79 A C ATOM 5337 N VAL 688 66.627 68.908 48.182 1.00 18.43 A N ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C										Α	С .	
ATOM 5329 0 SER 686 63.447 71.611 47.128 1.00 21.27 A 0 ATOM 5330 N THR 687 65.362 72.307 48.060 1.00 19.02 A N ATOM 5331 CA THR 687 66.122 71.189 47.509 1.00 17.15 A C ATOM 5332 CB THR 687 67.441 71.665 46.906 1.00 16.10 A C ATOM 5333 0G1 THR 687 68.362 71.959 47.960 1.00 17.42 A 0 ATOM 5334 CG2 THR 687 67.214 72.920 46.058 1.00 14.71 A C ATOM 5335 C THR 687 66.433 70.153 48.585 1.00 15.79 A C ATOM 5336 0 THR 687 66.496 70.466 49.763 1.00 15.82 A 0 ATOM 5337 N VAL 688 66.627 68.908 48.182 1.00 18.43 A N ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C	ATOM									Α		
ATOM 5330 N THR 687 65.362 72.307 48.060 1.00 19.02 A N ATOM 5331 CA THR 687 66.122 71.189 47.509 1.00 17.15 A C ATOM 5332 CB THR 687 67.441 71.665 46.906 1.00 16.10 A C ATOM 5333 0G1 THR 687 68.362 71.959 47.960 1.00 17.42 A O ATOM 5334 CG2 THR 687 67.214 72.920 46.058 1.00 14.71 A C ATOM 5335 C THR 687 66.433 70.153 48.585 1.00 15.79 A C ATOM 5336 O THR 687 66.496 70.466 49.763 1.00 15.82 A O ATOM 5337 N VAL 688 66.627 68.908 48.182 1.00 18.43 A N ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C										Α		
ATOM 5331 CA THR 687 66.122 71.189 47.509 1.00 17.15 A C ATOM 5332 CB THR 687 67.441 71.665 46.906 1.00 16.10 A C ATOM 5334 CG2 THR 687 67.214 72.920 46.058 1.00 14.71 A C ATOM 5335 C THR 687 66.433 70.153 48.585 1.00 14.71 A C ATOM 5336 O THR 687 66.496 70.466 49.763 1.00 15.79 A C ATOM 5337 N VAL 688 66.627 68.908 48.182 1.00 18.43 A N ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C												
ATOM 5332 CB THR 687 67.441 71.665 46.906 1.00 16.10 A C ATOM 5333 0G1 THR 687 68.362 71.959 47.960 1.00 17.42 A 0 ATOM 5334 CG2 THR 687 67.214 72.920 46.058 1.00 14.71 A C ATOM 5335 C THR 687 66.433 70.153 48.585 1.00 15.79 A C ATOM 5336 0 THR 687 66.496 70.466 49.763 1.00 15.82 A 0 ATOM 5337 N VAL 688 66.627 68.908 48.182 1.00 18.43 A N ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C												
ATOM 5333 OG1 THR 687 68.362 71.959 47.960 1.00 17.42 A O ATOM 5334 CG2 THR 687 66.433 70.153 48.585 1.00 14.71 A C ATOM 5336 O THR 687 66.496 70.466 49.763 1.00 15.79 A C ATOM 5337 N VAL 688 66.627 68.908 48.182 1.00 18.43 A N ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C												
ATOM 5334 CG2 THR 687 67. 214 72. 920 46. 058 1. 00 14. 71 A C ATOM 5335 C THR 687 66. 433 70. 153 48. 585 1. 00 15. 79 A C ATOM 5336 O THR 687 66. 496 70. 466 49. 763 1. 00 15. 82 A O ATOM 5337 N VAL 688 66. 627 68. 908 48. 182 1. 00 18. 43 A N ATOM 5338 CA VAL 688 66. 935 67. 854 49. 147 1. 00 17. 92 A C ATOM 5339 CB VAL 688 66. 840 66. 453 48. 480 1. 00 17. 13 A C												
ATOM 5335 C THR 687 66.433 70.153 48.585 1.00 15.79 A C ATOM 5336 0 THR 687 66.496 70.466 49.763 1.00 15.82 A O ATOM 5337 N VAL 688 66.627 68.908 48.182 1.00 18.43 A N ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C												
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ATOM 5337 N VAL 688 66.627 68.908 48.182 1.00 18.43 A N ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C												
ATOM 5338 CA VAL 688 66.935 67.854 49.147 1.00 17.92 A C ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C												
ATOM 5339 CB VAL 688 66.840 66.453 48.480 1.00 17.13 A C												
1 MOV 5010 001 1111 000												
	ATOM	5340	CG1	VAL	688					_	č	

					· FI	G. 4·	- 1 1 0			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5341 5342 5343 5344 5345 5346 5347 5348 5349 5350	C O N CA CB CG SD CE C	VAL VAL WET MET MET MET MET MET MET	688 689 689 689 689 689 689	65. 459 68. 341 68. 559 69. 280 70. 672 71. 475 71. 829 70. 465 70. 338 70. 897	66. 279 68. 059 67. 905 68. 428 68. 647 69. 213 69. 210 69. 539	50. 923 48. 851 49. 246 48. 065 46. 984 45. 909 44. 871 50. 479	1. 00 18. 49 1. 00 17. 50 1. 00 15. 69 1. 00 16. 92 1. 00 17. 40 1. 00 13. 91 1. 00 10. 55 1. 00 11. 73 1. 00 9. 36 1. 00 17. 90	A A A A A A A	CONTINUE OF CONTIN	,
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5351 5352 5353 5354 5355 5356 5357 5358 5359	O N CA CB OG C O N CA CB	MET SER SER SER SER SER ARG ARG	689 690 690 690 690 690 691 691	71. 721 70. 179 70. 358 69. 621 68. 234 69. 898 69. 930 69. 480 69. 041 67. 591	69. 220 70. 653 71. 544 72. 866 72. 702 70. 933 71. 606 69. 672 69. 012 68. 546	51. 341 50. 569 51. 712 51. 501 51. 711 53. 038 54. 063 53. 023 54. 249 54. 113	1. 00 16. 90 1. 00 18. 32 1. 00 21. 65 1. 00 20. 29 1. 00 24. 78 1. 00 22. 31 1. 00 23. 43 1. 00 21. 70 1. 00 23. 07 1. 00 22. 90	A A A A A A	O N C C O C O N C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5361 5362 5363 5364 5365 5366 5367 5368 5369 5370		ARG ARG ARG ARG ARG ARG ALA ALA	691 691 691 691 691 691 691 692 692	66. 623 65. 201 64. 236 62. 963 62. 509 62. 149 69. 922 69. 595 71. 041	69. 652 69. 152 70. 240 70. 134 68. 989 71. 172 67. 811 67. 635	53. 770 53. 813 53. 694 54. 061 54. 566 53. 946 54. 593 55. 488 53. 889	1. 00 22. 81 1. 00 22. 97 1. 00 24. 03 1. 00 26. 18 1. 00 25. 20 1. 00 26. 01 1. 00 24. 24 1. 00 25. 28 1. 00 24. 03	A A A A A A	C C N C N C O N	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5371 5372 5373 5374 5375 5376 5377 5378 5379	CB C O N CA CB CG	ALA ALA ALA GLU GLU GLU GLU GLU GLU	692 692 692 693 693 693 693 693	71. 960 73. 270 72. 251 72. 066 72. 707 73. 033 73. 351 74. 829 75. 604 74. 984	66. 561 66. 826 66. 210 65. 068 67. 181 66. 944 68. 266 68. 606 67. 627 66. 948	54. 100 53. 360 55. 562 55. 967 56. 347 57. 757 58. 463 58. 583 59. 463 60. 316	1. 00 24. 84 1. 00 24. 20 1. 00 24. 60 1. 00 25. 74 1. 00 27. 13 1. 00 29. 38 1. 00 35. 02 1. 00 39. 06 1. 00 38. 42	A A A A A A A A	C C O N C C C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5380 5381 5382 5383 5384 5385 5386 5387 5388 5389	OE2 C O N CA CB CG OD1 ND2 C	GLU GLU ASN ASN ASN ASN ASN	693 693 693 694 694 694 694 694 694	76. 845 71. 947 72. 250 70. 688 69. 594 68. 274 68. 191 67. 291 69. 127 69. 412	67. 554 66. 215 65. 505 66. 387 65. 734 66. 473 67. 796 68. 607 68. 015 64. 252	59. 307 58. 549 59. 506 58. 160 58. 873 58. 619 59. 370 59. 132 60. 287	1.00 41.03 1.00 26.16 1.00 26.78 1.00 24.46 1.00 24.35 1.00 26.79 1.00 28.23 1.00 29.60 1.00 27.09 1.00 22.78	A A A A A A A A	0 C O N C C C C	

·	•.			FIG. 4-112	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5444 5444 5444 5444 5444 5444 5444 5444 5444 5445 5445 5445 5446 5465	CG TYR CD1 TYR CD1 TYR CD2 TYR CC2 TYR C LEU C L	700 700 700 700 700 700 700 701 701 701	68. 374 58. 105 52. 114 1. 00 13. 03 A 67. 027 58. 171 51. 746 1. 00 12. 78 A 66. 611 58. 961 50. 666 1. 00 7. 94 A 69. 301 58. 840 51. 359 1. 00 12. 91 A 68. 895 59. 629 50. 282 1. 00 10. 45 A 67. 550 59. 688 49. 948 1. 00 10. 05 A 67. 150 60. 495 48. 913 1. 00 8. 37 A 68. 743 55. 056 52. 468 1. 00 11. 71 A 69. 881 54. 594 52. 463 1. 00 10. 84 A 67. 836 54. 775 51. 540 1. 00 11. 32 A 68. 142 53. 950 50. 383 1. 00 11. 03 A 67. 313 52. 667 50. 378 1. 00 8. 96 A 67. 439 51. 794 49. 123 1. 00 10. 04 A 68. 841 51. 873 48. 511 1. 00 7. 25 A 67. 089 50. 376 49. 490 1. 00 5. 44 A 67. 811 54. 799 49. 170 1. 00 13. 03 A 66. 660 55. 219 48. 986 1. 00 12. 91 A 68. 724 55. 888 47. 169 1. 00 11. 74 A 69. 806 56. 968 47. 196 1. 00 11. 17 A 69. 916 57. 965 46. 044 1. 00 12. 13 A 68. 569 58. 656 45. 803 1. 00 10. 37 A 68. 883 55. 003 45. 942 1. 00 13. 49 A 69. 854 54. 251 45. 832 1. 00 14. 04 A 67. 935 55. 111 45. 016 1. 00 13. 82 A 67. 934 54. 297 43. 806 1. 00 12. 92 A 66. 897 52. 305 42. 706 1. 00 12. 98 A 66. 897 52. 305 42. 706 1. 00 13. 52 A 66. 202 51. 383 45. 663 1. 00 13. 28 A 67. 561 55. 125 42. 582 1. 00 14. 12 A	CCCCCONCCCCCONCCCCONCCCCC
ATOM ATOM ATOM ATOM ATOM ATOM	5472 5473 5474 5475 5476 5477	N HIS CA HIS CB HIS CG HIS CD2 HIS ND1 HIS	704 704 704 704 704 704 704	68. 265 54. 909 41. 473 1. 00 13. 28 A 67. 987 55. 678 40. 265 1. 00 11. 81 A 68. 670 57. 048 40. 391 1. 00 11. 13 A 67. 968 58. 156 39. 667 1. 00 11. 66 A 67. 446 58. 221 38. 418 1. 00 10. 83 A	O N C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5478 5479 5480 5481 5482 5483 5484 5485 5486 5487	CE1 HIS NE2 HIS C HIS O HIS N GLY CA GLY C GLY O GLY N THR CA THR	704 704 704 704 705 705 705 705 706 706	67. 736 59. 387 40. 244 1. 00 10. 07 A 67. 098 60. 162 39. 385 1. 00 9. 04 A 66. 910 59. 479 38. 270 1. 00 11. 23 A 68. 464 54. 965 38. 992 1. 00 11. 87 A 69. 503 54. 306 38. 980 1. 00 11. 87 A 67. 684 55. 082 37. 926 1. 00 11. 49 A 68. 075 54. 486 36. 663 1. 00 11. 90 A 69. 066 55. 449 36. 036 1. 00 12. 16 A 68. 911 56. 660 36. 153 1. 00 13. 94 A 70. 086 54. 928 35. 372 1. 00 13. 29 A 71. 101 55. 782 34. 770 1. 00 12. 51	N C N C O N C C O N

			FIG. 4-113	(Continued)
ATOM	5488 CB TH	R 706		4 0
ATOM				A C
ATOM			=0 -1	A 0
ATOM			70 0-1	A C
ATOM	5492 0 TH			A C
ATOM	5493 N AL		71.183 57.461 33.084 1.00 14.35	A 0
ATOM	5494 CA ALA		69. 754 55. 770 32. 748 1. 00 13. 82	A N
ATOM	5495 CB ALA		69. 289 56. 302 31. 469 1. 00 15. 26	A C
ATOM	5496 C AL		69. 126 55. 176 30. 442 1. 00 13. 60	A C
ATOM			67. 970 57. 030 31. 644 1. 00 16. 56	A C
ATOM			67.154 57.075 30.720 1.00 17.71	A 0
ATOM			67.764 57.600 32.828 1.00 16.33	A N
ATOM	5499 CA ASF 5500 CB ASF		66. 534 58. 314 33. 113 1. 00 16. 71	A C
ATOM			66. 376 58. 508 34. 614 1. 00 18. 25	A C
ATOM			64.957 58.834 35.000 1.00 19.59	A C
ATOM			64. 304 59. 612 34. 266 1. 00 18. 82	A 0
ATOM			64.498 58.317 36.038 1.00 19.68	A 0
ATOM	•		66. 490 59. 673 32. 408 1. 00 17. 30	A C
ATOM	5505 0 ASP 5506 N ASP		67. 131 60. 647 32. 843 1. 00 18. 75	A 0
ATOM	5507 CA ASP		65. 715 59. 722 31. 327 1. 00 13. 98	A N
ATOM	5508 CB ASP			A C
ATOM	5509 CG ASP			A C
ATOM	5510 OD1 ASP	709 709		A C
ATOM	5511 OD2 ASP	709		A 0
ATOM	5512 C ASP	709		A 0
ATOM	5513 0 ASP	709		A C
ATOM	5514 N ASN	710		A 0
ATOM	5515 CA ASN	710		A N
ATOM	5516 CB ASN	710		A C
ATOM	5517 CG ASN	710		A C A C
ATOM	5518 OD1 ASN	710	EO 971 01 700 00 071	A C
ATOM	5519 ND2 ASN	710	60 691 69 590 99 999 1 99 1	A 0
ATOM	5520 C ASN	710	62 205 62 010 00 000	
ATOM	5521 0 ASN	710	69 601 64 911 99 919	
ATOM	5522 N VAL	711	62 570 69 946 95 017 1 00 11 40	-
ATOM	5523 CA VAL	711 -	63.570 62.246 35.017 1.00 11.10 A 64.221 62.741 36.225 1.00 9.96	
ATOM	5524 CB VAL	711	62 620 62 100 07 510	
ATOM	5525 CG1 VAL	711	64 415 69 570 00 710 4 00 5 00	
ATOM	5526 CG2 VAL	711	69 176 69 567 97 975	
ATOM	5527 C VAL	711	GE CAE CO ODE OG COO I CO	
ATOM	5528 0 VAL	711	6E 040 61 060 00 000	
ATOM	5529 N HIS	712	66 E10 C0 10C DE E01	
ATOM	5530 CA HIS	712	1,00 10.01	
ATOM	5531 CB HIS	712	100 1111	
ATOM	5532 CG HIS	712	100 100 1	
ATOM	5533 CD2 HIS	712	100 11.00	
ATOM	5534 ND1 HIS	712	1 100 12,00	
ATOM	5535 CE1 HIS	712	00 000 00 000	
ATOM	5536 NE2 HIS	712	0.0 0.0 0.1 0.11	
	2200 1110	114	66. 359 64. 944 31. 891 1. 00 11. 62 A	N

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			FIG. 4-114	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5537 C HIS 5538 O HIS 5539 N PHE 5540 CA PHE 5541 CB PHE 5542 CG PHE 5543 CD1 PHE 5544 CD2 PHE 5545 CE1 PHE 5546 CE2 PHE 5547 CZ PHE 5548 C PHE 5549 O PHE 5549 O PHE 5550 N GLN 5551 CA GLN 5551 CA GLN 5552 CB GLN 5553 CG GLN 5554 CD GLN 5555 OE1 GLN 5555 CG GLN 5556 NE2 GLN 5557 C GLN 5556 NE2 GLN 5556 NE2 GLN 5556 CA GLN 5561 CB GLN 5556 NE2 GLN 5557 C GLN 5560 CA GLN 5561 CB GLN 5561 CB GLN 5562 CG GLN 5563 CD GLN 5564 OE1 GLN 5565 NE2 GLN 5565 CA GLN 5567 O GLN 5567 O GLN 5568 N SER 5569 CA SER 5570 CB SER 5571 OG SER 5572 C SER 5573 O SER 5574 N ALA 5575 CA ALA	712 713 713 713 713 713 713 713 713 714 714 714 714 714 715 715 715 715 715 716 716 716 716 717	FIG. 4 - 1.14 68. 698 62. 222 36. 491 1. 00 10. 63 68. 461 62. 598 37. 633 1. 00 11. 98 69. 631 61. 319 36. 210 1. 00 10. 82 70. 458 60. 720 37. 251 1. 00 11. 00 71. 533 59. 823 36. 634 1. 00 11. 14 72. 270 58. 989 37. 639 1. 00 11. 47 71. 714 57. 813 38. 126 1. 00 11. 22 73. 496 59. 407 38. 144 1. 00 11. 84 72. 367 57. 066 39. 109 1. 00 11. 98 74. 153 58. 667 39. 126 1. 00 13. 82 73. 586 57. 495 39. 610 1. 00 11. 04 71. 122 61. 818 38. 061 1. 00 11. 85 71. 404 61. 640 39. 243 1. 00 13. 14 71. 377 62. 948 37. 403 1. 00 12. 47 72. 001 64. 113 38. 022 1. 00 10. 55 71. 851 65. 321 37. 082 1. 00 11. 91 72. 055 66. 695 37. 740 1. 00 10. 69 71. 501 67. 827 36. 891 1. 00 9. 77 70. 447 67. 693 36. 268 1. 00 10. 50 72. 201 68. 948 36. 870 1. 00 9. 43 71. 355 64. 417 39. 368 1. 00 9. 91 72. 037 64. 700 40. 356 1. 00 8. 86 70. 029 64. 340 39. 395 1. 00 10. 27 69. 255 64. 616 40. 599 1. 00 10. 62 67. 771 64. 393 40. 315 1. 00 10. 98 67. 267 65. 219 39. 144 1. 00 11. 10 66. 285 66. 288 39. 567 1. 00 14. 59 66. 381 66. 828 40. 671 1. 00 10. 98 67. 267 65. 219 39. 144 1. 00 11. 10 66. 285 66. 288 39. 567 1. 00 12. 90 69. 716 63. 781 41. 780 1. 00 10. 62 67. 771 64. 393 40. 315 1. 00 10. 98 69. 976 64. 322 42. 853 1. 00 12. 32 69. 828 62. 472 41. 600 1. 00 9. 91 70. 299 61. 630 42. 700 1. 00 12. 35 69. 937 60. 163 42. 461 1. 00 10. 77 68. 541 59. 994 42. 492 1. 00 14. 60 71. 818 61. 761 42. 876 1. 00 13. 46 72. 3241 61. 556 43. 976 1. 00 14. 90 72. 522 62. 094 41. 797 1. 00 12. 22	
ATOM	5576 CB ALA	717 717	73. 969 62. 252 41. 870 1. 00 13. 92 A 74. 555 62. 487 40. 479 1. 00 12. 46 A	C C
ATOM ATOM ATOM	5577 C ALA 5578 O ALA 5579 N GLN	717 717	74. 299 63. 423 42. 790 1. 00 13. 73 A 75. 257 63. 375 43. 560 1. 00 15. 24 A	Č O
ATOM	5580 CA GLN	718 718	73. 504 64. 482 42. 710 1. 00 13. 27 73. 738 65. 631 43. 565 1. 00 13. 07	N C
ATOM ATOM	5581 CB GLN 5582 CG GLN	718 718	72. 976 66. 841 43. 035 1. 00 13. 93 A	C
ATOM	5583 CD GLN	718	73. 548 67. 422 41. 734 1. 00 15. 44 A 74. 996 67. 865 41. 867 1. 00 13. 84 A	C C
ATOM ATOM	5584 OE1 GLN 5585 NE2 GLN	718	75. 467 68. 172 42. 950 1. 00 16. 85 A	0
ATOM	5585 NE2 GLN	718	75. 699 67. 915 40. 755 1. 00 17. 86 A	N

		FIG. 4-115	(Continued)
ATON ATON	5587 0 GLN 718	73. 350 65. 343 45. 026 1. 00 13. 24 73. 941 65. 910 45. 949 1. 00 11. 74	A C
ATON	1 5588 N ILE 719	72. 370 64. 460 45. 237 1. 00 12. 01	A O
ATOM	170	71. 956 64. 110 46. 594 1. 00 11. 94	A N A C
ATOM	110	70. 691 63. 201 46. 616 1. 00 12. 50	A C A C
ATOM	110	70. 464 62. 673 48. 021 1. 00 11. 09	A C
ATOM	110	69.447 63.979 46.174 1.00 14.37	A Č
ATOM	110	68. 170 63. 143 46. 154 1. 00 8 64	A Č
ATOM ATOM	122 110	73. 081 63. 338 47. 282 1. 00 11. 72	Ä Č
ATOM		73. 543 63. 703 48. 367 1. 00 10. 69	A O
ATOM	120	73. 508 62. 262 46. 632 1. 00 11. 35	A N
ATOM	5597 CA SER 720 5598 CB SER 720	74. 557 61. 405 47. 155 1. 00 11. 02	A C
ATOM	5599 OG SER 720	74. 901 60. 325 46. 135 1. 00 10. 89	A C
ATOM	5600 C SER 720	75. 471 60. 894 44. 970 1. 00 13. 75 75. 804 62. 207 47. 488 1. 00 12. 63	A 0
ATOM	5601 0 SER 720	70 100	A C
ATOM	5602 N LYS 721	70 470	A 0
ATOM	5603 CA LYS 721	77 000	A N
ATOM	5604 CB LYS 721	77. 336 63. 951 46. 800 1. 00 12. 15 77. 613 64. 823 45. 571 1. 00 11. 24	A C
ATOM	5605 CG LYS 721	78. 764 65. 796 45. 756 1. 00 7. 41	A C
ATOM	5606 CD LYS 721	79. 517 66. 064 44. 451 1. 00 9. 30	A C A C
ATOM	5607 CE LYS 721	78. 674 66. 765 43. 392 1. 00 8. 74	A C A C
ATOM ATOM	5608 NZ LYS 721	78. 341 68. 165 43. 739 1. 00 9. 54	A N
ATOM	5609 C LYS 721 5610 O LYS 721	77. 190 64. 816 48. 038 1. 00 13. 24	Ä Č
ATOM		78. 150 64. 982 48. 791 1. 00 14. 49	A 0
ATOM	5611 N ALA 722 5612 CA ALA 722	75. 992 65. 351 48. 262 1. 00 13. 05	A N
ATOM	5613 CB ALA 722	75. 760 66. 198 49. 432 1. 00 13. 21	A C
ATOM	5614 C ALA 722	74. 389 66. 870 49. 353 1. 00 9. 37 75. 874 65. 369 50. 702 1. 00 14 04	A C
ATOM	5615 O ALA 722	70 100 11,01	A C
ATOM	5616 N LEU 723	75 000	A 0
ATOM	5617 CA LEU 723	75. 360 64. 145 50. 665 1. 00 14. 96 75. 429 63. 266 51. 826 1. 00 17. 23	A N
ATOM	5618 CB LEU 723	74. 626 61. 984 51. 570 1. 00 16. 86	A C
ATOM	5619 CG LEU 723	73. 116 62. 205 51. 463 1. 00 18. 78	A C A C
ATOM	5620 CD1 LEU 723	72. 428 60. 932 50. 991 1. 00 18. 74	
ATOM	5621 CD2 LEU 723	72. 576 62. 663 52. 817 1. 00 16. 86	A C .
ATOM ATOM	5622 C LEU 723	76. 889 62. 926 52. 134 1. 00 17. 26	A C A C
ATOM	5623 0 LEU 723 5624 N VAL 724	77. 320 62. 990 53. 280 1. 00 18. 48	A O
ATOM		77.641 62.559 51.103 1.00 17.41	A N
ATOM	5625 CA VAL 724 5626 CB VAL 724	79. 050 62. 234 51. 257 1. 00 16. 64	A C
ATOM	5627 CG1 VAL 724	79. 671 61. 824 49. 902 1. 00 14. 31	A C
ATOM	5628 CG2 VAL 724	81. 187 61. 819 49. 987 1. 00 13. 56 79. 178 60. 449 49. 519 1. 00 14. 78	A C
ATOM	5629 C VAL 724	50 70 11.10	A C
ATOM	5630 0 VAL 724	00 000	A C
ATOM	5631 N ASP 725	70 111	A 0
ATOM	5632 CA ASP 725	79. 411 64. 632 51. 318 1. 00 19. 19 80. 051 65. 848 51. 776 1. 00 20. 26	A N
ATOM	5633 CB ASP 725	79. 627 67. 032 50. 919 1. 00 22. 40	A C A C
ATOM	5634 CG ASP 725	80. 259 67. 004 49. 549 1. 00 26. 44	A C A C
			·• v

					FIC	3.4-	116			(Conti	nued)
ATOM ATOM ATOM	5635 5636 5637	OD1 OD2 C		725 725 725	81.149 79.867 79.805	66. 151 67. 839 66. 171	49. 319 48. 704 53. 238	1.00 26.28 1.00 30.70 1.00 19.86	A A A	0 0 C	
ATOM ATOM	5638 5639	0 N	ASP VAL	725 726	80. 486 78. 841	67. 024 65. 516	53. 792 53. 873	1.00 23.33 1.00 17.95	A A	O N	
ATOM ATOM ATOM	5640 5641 5642		VAL VAL VAL	726 726 726	78.603 77.178 76.992	65. 790 66. 341 67. 680	55. 285 55. 567 54. 875	1.00 17.97 1.00 18.54 1.00 16.64	A A A	. C C	
ATOM ATOM ATOM	5643 5644 5645		VAL VAL VAL	726 726 726	76. 121 78. 812 78. 412	65. 339 64. 549 64. 504	55. 120 56. 124 57. 283	1.00 18.24 1.00 17.82 1.00 19.86	A A A	C C 0	
ATOM ATOM	5646 5647	N CA	GLY GLY	727 727	79. 439 79. 711 78. 509	63. 541 62. 317 61. 489	55. 535 56. 263 56. 681	1.00 17.13 1.00 16.84 1.00 17.94	. A . A	N C C	
ATOM ATOM ATOM	5648 5649 5650	0 N	GLY GLY VAL	727 727 728	78. 483 77. 517	60. 961 61. 371	57. 794 55. 802	1.00 19.74 1.00 16.62	A A	O N	
ATOM ATOM ATOM	5651 5652 5653	CA CB CG1	VAL VAL VAL	728 728 728	76. 331 75. 030 73. 838	60. 571 61. 302 60. 338	56.085 55.643 55.668	1.00 17.26 1.00 18.46 1.00 16.22	A A A	C C C	
ATOM ATOM ATOM	5654 5655 5656	CG2 C O	VAL VAL VAL	728 728 728	74. 753 76. 411 76. 667	62. 476 59. 230 59. 186	56. 579 55. 347 54. 143	1.00 18.70 1.00 18.03 1.00 18.40	A A A	C C 0	
ATOM ATOM ATOM	5657 5658 5659	N CA CB	ASP ASP ASP	729 729 729	76. 211 76. 246 76. 734	58. 135 56. 822 55. 752	56.069 55.441 56.420	1.00 18.22 1.00 19.90 1.00 22.57	A A A	· N C C	
ATOM ATOM	5660 5661	CG OD1	ASP ASP	729 729	76. 819 77. 340 76. 372	54. 376 54. 278 53. 388	55. 778 54. 649 56. 398	1.00 25.97 1.00 27.13 1.00 30.03	A A A	C 0 0	
ATOM ATOM ATOM	5662 5663 5664	OD2 C O	ASP ASP	729 729 729	74. 839 73. 868	56. 504 56. 863	54.984 55.649	1.00 19.16 1.00 21.91	A A	C 0	
ATOM ATOM ATOM	5665 5666 5667	N CA CB	PHE PHE PHE	730 730 730	74. 723 73. 416 72. 796	55. 838 55. 499 56. 734	53. 846 53. 299 52. 639	1.00 18.27 1.00 16.06 1.00 14.49	A A A	N C C	
ATOM ATOM ATOM	5668 5669 5670		PHE PHE PHE	730 730 730	73. 590 73. 262 74. 691	57. 265 56. 913 58. 082	51. 480 50. 177 51. 694	1.00 12.02 1.00 10.26 1.00 11.55	A A A	C C C	
ATOM ATOM ATOM	5671 5672 5673	CE1	PHE PHE PHE	730 730 730	74. 020 75. 459 75. 120	57. 364 58. 537 58. 175	49. 098 50. 621 49. 317	1.00 10.41 1.00 13.40 1.00 9.85	A A A	C C C C	
ATOM ATOM	5674 5675	C 0	PHE PHE	730 730	73. 565 74. 675	54. 388 53. 990	52. 281 51. 945	1.00 16.20 1.00 18.49	A A	0	
ATOM ATOM ATOM	5676 5677 5678	N CA CB	GLN GLN GLN	731 731 731	72. 447 72. 484 71. 514	53. 883 52. 813 51. 708	51. 791 50. 813 51. 208	1.00 17.40 1.00 17.82 1.00 20.04	A A A	N C C	
ATOM ATOM ATOM	5679 5680 5681	CG CD OE1	GLN GLN GLN	731 731 731	71. 641 73. 019 73. 554	51. 257 50. 737 49. 883	52. 644 52. 968 52. 256	1.00 25.37 1.00 28.25 1.00 32.85	A A A	C C 0	
ATOM ATOM	5682 5683	NE2 C	GLN GLN	731 731	73. 603 72. 091	51. 238 53. 382	54. 055 49. 458	1.00 30.12 1.00 17.65	A A	N C	

	FIG. 4-117	(Continued)
ATOM 5684 O GLN 731 ATOM 5685 N ALA 732 ATOM 5686 CA ALA 732 ATOM 5687 CB ALA 732 ATOM 5688 C ALA 732 ATOM 5689 O ALA 732 ATOM 5690 N MET 733 ATOM 5691 CA MET 733 ATOM 5692 CB MET 733 ATOM 5693 CG MET 733 ATOM 5694 SD MET 733 ATOM 5695 CE MET 733 ATOM 5696 C MET 733 ATOM 5696 C MET 733 ATOM 5697 O MET 733 ATOM 5698 N TRP 734 ATOM 5699 CA TRP 734 ATOM 5699 CA TRP 734 ATOM 5700 CB TRP 734 ATOM 5701 CG TRP 734 ATOM 5702 CD2 TRP 734 ATOM 5704 CE3 TRP 734 ATOM 5705 CD1 TRP 734 ATOM 5706 NE1 TRP 734 ATOM 5706 NE1 TRP 734 ATOM 5707 CZ2 TRP 734 ATOM 5708 CZ3 TRP 734 ATOM 5709 CH2 TRP 734 ATOM 5709 CH2 TRP 734 ATOM 5709 CH2 TRP 734 ATOM 5710 C TRP 734 ATOM 5711 C TRP 734 ATOM 5709 CH2 TRP 734 ATOM 5710 C TRP 734 ATOM 5711 C TRP 734 ATOM 5711 C TRP 734 ATOM 5712 N TYR 735 ATOM 5713 CA TYR 735 ATOM 5714 CB TYR 735 ATOM 5715 CG TYR 735 ATOM 5716 CD1 TYR 735 ATOM 5717 CE1 TYR 735 ATOM 5718 CD2 TYR 735 ATOM 5719 CE2 TYR 735 ATOM 5720 CZ TYR 735 ATOM 5721 OH TYR 735 ATOM 5721 OH TYR 735 ATOM 5722 C TYR 735 ATOM 5723 O TYR 735 ATOM 5724 N THR 736 ATOM 5725 CA THR 736 ATOM 5726 CB THR 736 ATOM 5727 OG1 THR 736 ATOM 5728 CG2 THR 736 ATOM 5729 C THR 736 ATOM 5730 O THR 736 ATOM 5731 N ASP 737	FIG. 4 - 1 1 7 71. 160 54. 191 49. 355 1. 00 17. 02 72. 802 52. 962 48. 421 1. 00 14. 78 72. 510 53. 444 47. 088 1. 00 15. 21 73. 588 54. 409 46. 626 1. 00 15. 17 72. 419 52. 282 46. 131 1. 00 15. 21 72. 940 51. 207 46. 396 1. 00 16. 17 71. 737 52. 504 45. 019 1. 00 14. 57 71. 599 51. 483 44. 008 1. 00 14. 86 70. 490 50. 499 44. 383 1. 00 15. 14 70. 288 49. 386 43. 353 1. 00 18. 04 71. 814 48. 476 42. 961 1. 00 22. 04 71. 892 47. 307 44. 310 1. 00 17. 75 71. 283 52. 153 42. 683 1. 00 14. 93 70. 317 52. 915 42. 574 1. 00 13. 82 71. 890 52. 447 40. 356 1. 00 13. 13 73. 173 53. 117 39. 827 1. 00 10. 39 74. 187 52. 159 39. 267 1. 00 8. 77 75. 398 51. 726 39. 894 1. 00 7. 74 75. 984 50. 757 39. 053 1. 00 9. 97 76. 045 52. 062 41. 087 1. 00 12. 87 77. 183 50. 119 39. 369 1. 00 9. 94 77. 238 51. 428 41. 400 1. 00 9. 32 77. 793 50. 468 40. 545 1. 00 13. 91 77. 183 50. 119 39. 369 1. 00 9. 94 77. 238 51. 428 41. 400 1. 00 9. 32 77. 793 50. 468 40. 545 1. 00 13. 91 70. 635 51. 570 38. 461 1. 00 14. 06 71. 903 50. 155 39. 653 1. 00 19. 94 71. 480 51. 291 39. 445 1. 00 14. 06 71. 903 50. 155 39. 653 1. 00 19. 94 77. 238 51. 570 38. 461 1. 00 15. 51 68. 705 50. 326 37. 556 1. 00 14. 96 68. 619 48. 081 38. 708 1. 00 14. 76 68. 619 48. 081 38. 708 1. 00 14. 45 68. 360 47. 278 39. 816 1. 00 14. 33 67. 696 49. 971 39. 848 1. 00 15. 54 67. 772 47. 835 40. 938 1. 00 14. 45 68. 360 47. 278 39. 816 1. 00 14. 52 67. 772 47. 835 40. 938 1. 00 15. 54 70. 625 50. 966 36. 104 1. 00 15. 52 71. 763 50. 330 35. 654 1. 00 15. 54 72. 361 50. 608 34. 353 1. 00 15. 13 73. 491 49. 602 34. 030 1. 00 14. 68 74. 470 49. 614 35. 076 1. 00 15. 44 72. 361 50. 608 34. 353 1. 00 15. 44 72. 361 50. 608 34. 353 1. 00 15. 44 72. 361 50. 608 34. 353 1. 00 15. 44 72. 361 50. 608 34. 353 1. 00 15. 44 72. 361 50. 608 34. 353 1. 00 15. 44 72. 361 50. 608 34. 353 1. 00 15. 44 72. 361 50. 608 34. 353 1. 00 15. 44 72. 361 50. 608 34. 353 1. 00 15. 44 72. 361 50. 608 34. 353 1. 00 15. 44 74. 156 49. 961 32. 713 1. 00 14. 75 71. 365 50. 549 33. 206 1. 00 15. 44 71	A O A N A C A A A C A A A C A A A C A A A C A A A C A A A C A A A C A A A C A A A C A A A

										(Cont	inued)
					FIC	G. 4 -	118			(COIII)	mucu
ATOM	5733	СВ	ASP	7 37	70. 884	50. 677	30. 200	1.00 15.90	A	C	
ATOM	5734	CG	ASP	737	72. 232	50. 972	29. 574	1.00 13.30	A	C C	
ATOM	5735		ASP		72. 679	50. 147	28. 747	1.00 20.37	A		
ATOM	5736		ASP	737	72. 847	52. 020	29. 895	1.00 24.29	A	0 0	
ATOM	5737	C	ASP	737	68. 974	51.632	31. 467	1.00 18.74	A A	C	
ATOM	5738	Ö	ASP	737	68. 205	51.507	30. 515	1.00 18.86	A	Ö	
ATOM	5739	N	GLU	738	68. 553	51.692	32. 722	1.00 18.39	A	N	
ATOM	5740	CA	GLU	738	67. 135	51.644	33. 033	1.00 19.00	A	C	
ATOM	5741	CB	GLU	738	66. 909	50. 999	34. 407	1.00 20.24	A	C	•
ATOM	5742	CG	GLU	738	66. 904	49.485	34. 380	1.00 20.24	A	Č	
ATOM	5743	CD	GLU	738	65. 741	48. 937	33. 565	1.00 24.58	A	Č	
ATOM	5744	0E1		738	64. 588	49. 289	33. 878	1.00 27.21	Ä	0	
ATOM	5745		GLU	738	65.970	48. 163	32.611	1.00 26.16	A	ő	
ATOM	5746	C	GLU	738	66.624	53.076	33.025	1.00 19.38	Ä	Č.	
ATOM	5747	0	GLU	738	67.327	53.991	33.461	1.00 20.83	A	Ö	
ATOM	5748	N	ASP	739	65.414	53. 288	32.525	1.00 18.55	Ā	N	
ATOM	5749	CA	ASP	739	64.892	54.642	32.493	1.00 17.49	Α	C	
ATOM	5750	CB	ASP	739	64.074	54.863	31.222	1.00 18.32	Α	C	
ATOM	5751	CG	ASP	739	62.689	54:271		1.00 21.44	Α	С	
ATOM	5752		ASP	739	61.995	54.340	30. 257	1.00 24.73	Α	0	
ATOM	5753		ASP	739	62. 285	53. 752	32.358	1.00 21.35	Α	0	
ATOM	5754	C	ASP	739	64.088	54.976	33. 750	1.00 17.35	Α	C	
ATOM	5755	0	ASP	739	64. 191	54. 282	34. 762	1.00 15.74	A	0	
ATOM	5756	N	HIS	740	63. 291	56. 034	33. 687	1.00 16.96	A	N	
ATOM ATOM	5757	CA	HIS	740	62. 521	56.469	34. 842	1.00 18.24	A	C	
ATOM	5758 5759	CB CG	HIS HIS	740	61.746	57. 736	34.511	1.00 16.88	A	C	
ATOM	5760		HIS	740 740	61.145	58. 392	35. 710,	1.00 17.57	A	C	
ATOM	5761		HIS	740	59. 883 61. 881	58. 812 58. 687	35.961	1.00 16.26	A	C	
ATOM	5762		HIS	740	61.097	59. 262	36. 837 37. 732	1.00 17.31	A	N	
ATOM	5763		HIS	740	59. 880	59. 349	37. 224	1.00 18.51 1.00 17.94	A	C	
ATOM	5764		HIS	740	61.557	55. 449	35. 426	1.00 17.94	A	N	
ATOM	5765	ŏ	HIS	740	61.191	55. 539	36. 599	1.00 19.30	A A	C 0	
ATOM	5766	Ň	GLY	741	61.151	54. 481	34.614	1.00 20.00	A	N N	
ATOM	5767	CA	GLY	741	60. 216	53. 484	35. 084	1.00 18.82	Ä	C	•
ATOM	5768	C	GLY	741	60. 849	52. 218	35.609	1.00 20.36	A	Č	
ATOM	5769	0	GLY	741	60.165	51.404	36. 237	1.00 22.79	A	ŏ	
ATOM	5770	N	ILE	742	62.145	52.045	35.368	1.00 19.61	Ä	Ň	
ATOM	5771	CA	ILE	742	62.854	50.849	35.821	1.00 17.74	Ä	Ċ	
ATOM	5772	CB	ILE	742	63.273	50.981	37.294	1.00 14.46	Ä	č	
ATOM	5773		ILE	742	64.279	49.917	37.638	1.00 14.37	Ā	Č	
ATOM	5774		ILE	742	63.865	52.370	37.540	1.00 13.43	Α	Ċ	
ATOM	5775		ILE	742	64.540	52. 552	38.887	1.00 9.55	Α	C	
ATOM	5776	C	ILE	742	61.907	49.658	35.676	1.00 19.11	Α	С	
ATOM	5777	0	ILE	742	61.805	48. 825	36.571	1.00 18.97	Α	0	
ATOM	5778	N	ALA	743	61.217	49. 594	34. 534	1.00 20.16	A	N	-
ATOM .	5779	CA	ALA	743	60. 246	48. 538	34. 268	1.00 19.71	A	C	
ATOM	5780 5781	CB	ALA	743	59.004	49. 141	33.630	1.00 19.65	A	C	
ATOM	5781	С	ALA	743	60.717	47.350	33.430	1.00 20.08	Α	С	

					•		(Continued)
				FIG. 4	- 119		(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5782 0 5783 N 5784 CA 5785 CE 5786 00 5787 C 5788 0 5789 N 5790 CA 5791 CB 5792 0G 5793 C	SER SER SER SER SER SER SER SER SER	743 744 744 744 744 745 745 745 745 745		33. 006 1. 00 20. 9 33. 163 1. 00 19. 1 32. 389 1. 00 17. 3 32. 068 1. 00 14. 6 33. 125 1. 00 18. 0 33. 300 1. 00 16. 5 34. 519 1. 00 15. 4 32. 715 1. 00 19. 16 33. 503 1. 00 20. 00 32. 598 1. 00 20. 15 33. 377 1. 00 27. 90	2 A 4 A 2 A 4 A 8 A 7 A 0 A 3 A 2 A	(Continued) 0 N C C 0 C 0 C 0 C 0 C 0 C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5799 CG 5800 C 5801 O	THR 1 THR 2 THR THR THR	745 746 746 746 746 746 746 746	62. 130 42. 078 63. 783 42. 158 64. 796 41. 849 66. 125 41. 538 66. 463 42. 615 66. 009 40. 259 64. 996 42. 966 65. 066 42. 706	35. 781	3 A A A A A A A	0 N C C O C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5802 N 5803 CA 5804 CB 5805 C 5806 O 5807 N 5808 CA 5809 CB	ALA ALA ALA ALA HIS HIS	747 747 747 747 747 748 748 748	65. 070 44. 208 65. 286 45. 334 65. 554 46. 609 64. 113 45. 540 64. 291 45. 989 62. 915 45. 206 61. 718 45. 342 60. 477 45. 005	35. 821 1. 00 18. 73 36. 723 1. 00 18. 03 35. 919 1. 00 15. 38 37. 681 1. 00 17. 35 38. 814 1. 00 18. 52 37. 224 1. 00 16. 75 38. 046 1. 00 16. 92 37. 220 1. 00 13. 48	A A A A A	N C C C O N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5812 ND1 5813 CE1	HIS HIS HIS HIS HIS HIS GLN	748 748 748 748 748 748 748 749	59. 214 44. 968 58. 397 43. 941 58. 663 46. 094 57. 561 45. 762 57. 377 44. 461 61. 790 44. 415 61. 525 44. 816 62. 148 43. 165 62. 241 42. 201	38. 020 1. 00 14. 10 38. 348 1. 00 12. 63 38. 595 1. 00 14. 71 39. 241 1. 00 13. 05 39. 107 1. 00 14. 46 39. 263 1. 00 18. 16 40. 394 1. 00 20. 72 39. 025 1. 00 18. 81 40. 105 1. 00 19. 53	A A A A A A	C C N C N C O
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5823 NE2 5824 C 5825 O 5826 N 5827 CA 5828 CB 5829 CG	GLN GLN GLN GLN GLN HIS HIS HIS HIS	749 749 749 749 749 749 749 750 750 750 750	62. 408	39. 519 1. 00 20. 05 38. 550 1. 00 21. 82 37. 757 1. 00 20. 87 38. 316 1. 00 22. 37 36. 447 1. 00 20. 00 41. 008 1. 00 19. 07 42. 231 1. 00 17. 88 40. 399 1. 00 18. 97 41. 160 1. 00 16. 68 40. 226 1. 00 14. 65 40. 889 1. 00 13. 97	A A A A A A A A	C C C C O N C O N C C C

					FIC	3. 4·	- 120) *		(Continued)
ATOM ATOM ATOM	5831 5832 5833 5834	CE NE C	1 HIS 1 HIS 2 HIS HIS	750 750 750 750	68. 615 69. 804 70. 185 65. 529	44. 365 44. 000 42. 927 44. 400	42.320 41.653 42.157	1.00 12.57 1.00 12.04 1.00 17.33	A A A	N C N C
ATOM ATOM ATOM ATOM	5835 5836 5837 5838	O N CA CB	HIS ILE ILE ILE	750 751 751 751	65. 945 64. 899 64. 704 64. 206	44. 277 45. 490 46. 632 47. 849	41.726 42.604	1.00 17.03 1.00 15.90	A A A	.0 N C C
ATOM ATOM ATOM ATOM	5839 5840 5841 5842	CG: CD: C	ILE ILE ILE ILE	751 751 751 751	62. 893 64. 065 63. 684 63. 751	47. 504 49. 058 50. 332 46. 341	42.736	1.00 15.94 1.00 12.51	A A A	C C C C
ATOM ATOM ATOM ATOM	5843 5844 5845 5846	O N CA CB	ILE TYR TYR TYR	751 752 752 752	64. 062 62. 596 61. 651 60. 323	46. 632 45. 759 45. 449 44. 967	44. 919 43. 480 44. 551 43. 968	1.00 16.37	A A A A	O N C C
ATOM ATOM ATOM ATOM	5847 5848 5849 5850	CD2	TYR TYR	752 752 752 752	59. 443 58. 840 58. 102 59. 279	46. 126 46. 899 48. 026 46. 510	43. 593 44. 580 44. 258 42. 260	1.00 12.59 1.00 11.61 1.00 9.67 1.00 12.75	A A A	C C C C
ATOM ATOM ATOM ATOM ATOM	5851 5852 5853 5854 5855	CZ OH C	TYR TYR TYR TYR	752 752 752 752	58. 543 57. 964 57. 278 62. 226	47. 644 48. 395 49. 542 44. 429	41. 930 42. 940 42. 642 45. 522	1.00 10.28 1.00 9.02 1.00 12.10 1.00 16.42	A A A	C C O C
ATOM ATOM ATOM ATOM	5856 5857 5858 5859	O N CA CB OG1	TYR THR THR THR THR	752 753 753 753	61. 927 63. 056 63. 700 64. 502	44. 467 43. 526 42. 521 41. 510	46. 719 45. 004 45. 835 44. 985	1.00 16.42 1.00 15.74 1.00 16.30 1.00 15.57	A A A	O N C C
ATOM ATOM ATOM ATOM	5860 5861 5862 5863		THR THR THR THR	753 753 753 753	64. 678 64. 788	40. 677 40. 641 43. 240 42. 923	44. 253 45. 870 46. 758 47. 941	1.00 15.74 1.00 10.01 1.00 18.17 1.00 19.02	A A A	0 C C
ATOM ATOM ATOM ATOM	5864 5865 5866 5867	CA CB CG	HIS HIS HIS HIS	754 754 754 754 754	66. 363 67. 189 68. 449	44. 215 44. 972 45. 857 46. 379	46. 199 46. 959 46. 023 46. 644	1.00 18.78 1.00 18.90 1.00 19.13 1.00 19.62	A A A	N C C C
ATOM ATOM ATOM ATOM	5868 5869 5870	ND1 CE1	HIS HIS HIS HIS	754 754 754 754	69. 539 70. 493 70. 062	47. 619 45. 576 46. 298 47. 541	47. 070 46. 904 47. 462 47. 574	1.00 18.70 1.00 18.44 1.00 17.52 1.00 19.51	A A A	C N C N
ATOM ATOM ATOM ATOM	5872 5873 5874	O N CA CB	HIS MET MET MET	754 755 755	66. 088 64. 589 63. 854	45. 828 45. 876 46. 502 47. 342	48. 007 49. 158 47. 615 48. 558	1.00 19.38 1.00 19.63 1.00 18.83 1.00 19.68	A A A	C O N C
ATOM ATOM ATOM ATOM	5876 5877 5878	CG SD CE	MET MET MET MET	755 755 755 755 755	63. 283 4 62. 016 4 61. 100 4	48. 136 49. 173 50. 314 49. 270 46. 506	47. 839 46. 876 46. 309 45. 200 49. 676	1.00 16.86 1.00 16.00 1.00 20.78 1.00 15.61 1.00 20.27	A A A A	C C S C

				FIC 4-191	(Continued)
ATOM	5880	0 MET	755	F I G. 4 - 1 2 1 63.112 46.969 50.811 1.00 20.56 A	0
ATOM	5881		756	63.112 46.969 50.811 1.00 20.56 A 62.842 45.276 49.352 1.00 20.59 A	O N
ATOM	5882		756	62.240 44.380 50.332 1.00 21.43 A	Ċ
ATOM	5883		756	61.740 43.106 49.646 1.00 21.74 A	Č
ATOM	5884		756	60. 598 43. 373 48. 850 1. 00 21. 68 A	0
ATOM ATOM	5885 5886		756	63. 224 44. 023 51. 444 1. 00 22. 50 A	C
ATOM	5887		756 757	62. 858 44. 022 . 52. 623 1. 00 22. 47 A 64. 466 43. 716 51. 073 1. 00 22. 47 A	0
ATOM	5888		757	0F 400 40 004 F0 00F	N
ATOM	5889		757	65. 483 43. 384 52. 065 1. 00 23. 01 A 66. 828 43. 032 51. 407 1. 00 21. 90 A	C C
ATOM	5890	CG HIS	757	66. 837 41. 721 50. 682 1. 00 24. 99 A	Č
ATOM	5891	CD2 HIS	757	67. 344 41. 375 49. 473 1. 00 26. 07 A	č
ATOM.	5892	ND1 HIS	757 757	66. 314 40. 563 51. 220 1. 00 26. 51 A	N
ATOM ATOM	5893 5894	CE1 HIS NE2 HIS	757 757	66. 497 39. 564 50. 375 1. 00 25. 15 A	C
ATOM	5895	C HIS	757	67.120 40.029 49.307 1.00 25.93 A 65.689 44.596 52.966 1.00 23.03 A	N
ATOM	5896	0 HIS	757	65. 689 44. 596 52. 966 1. 00 23. 03 A 65. 823 44. 474 54. 186 1. 00 24. 03 A	C 0
ATOM	5897	N PHE	758	65. 704 45. 771 52. 356 1. 00 22. 28 A	N N
ATOM	5898	CA PHE	758	65. 920 46. 995 53. 106 1. 00 24. 10 A	Č
ATOM ATOM	5899 5900	CB PHE	758 758	66.005 48.190 52.161 1.00 20.12 A	C
ATOM	5900	CG PHE CD1 PHE	758 758	66. 455 49. 448 52. 828 1. 00 17. 08 A	Ċ
ATOM	5902	CD2 PHE	758	67. 803 49. 657 53. 106 1. 00 15. 49 A 65. 537 50. 429 53. 176 1. 00 15. 44 A	C
ATOM	5903	CE1 PHE	758	65. 537 50. 429 53. 176 1. 00 15. 44 A 68. 233 50. 825 53. 717 1. 00 14. 07 A	C
ATOM	5904	CE2 PHE	758	65. 955 51. 607 53. 789 1. 00 17. 18 A	C
ATOM	5905	CZ PHE	758	67. 308 51. 806 54. 060 1. 00 15. 05 A	č
ATOM ATOM	5906	C PHE	758 758	64. 832 47. 254 54. 135 1. 00 26. 28 A	C C
ATOM	5907 5908	O PHE N ILE	758 759	65. 120 47. 546 55. 295 1. 00 28. 09 A	0
ATOM	5909	CA ILE	759	63. 580 47. 162 53. 706 1. 00 27. 69 A 62. 461 47. 394 54. 605 1. 00 29. 02 A	N
ATOM	5910	CB ILE	759	62. 461 47. 394 54. 605 1. 00 29. 02 A 61. 129 47. 271 53. 853 1. 00 28. 24 A	C C
ATOM	5911	CG2 ILE	759	59. 967 47. 207 54. 836 1. 00 29. 09 A	Č
ATOM	5912	CG1 ILE	759	60. 990 48. 446 52. 884 1. 00 28. 85 A	č
ATOM ATOM	5913	CD1 ILE	759	61.173 49.809 53.535 1.00 27.28 A	Ċ
ATOM	5914 5915	C ILE O ILE	759 759	62. 467 46. 420 55. 774 1. 00 31. 10 A	C
ATOM	5916	N LYS	760	62. 292 46. 822 56. 925 1. 00 30. 20 A 62. 669 45. 140 55. 464 1. 00 32. 71 A	0
ATOM	5917	CA LYS	760	69 607 44 070 56 407 4 60 60	N C
ATOM	5918	CB LYS	760	62. 732 42. 715 55. 780 1. 00 34. 00 A	C C
ATOM	5919	CG LYS	760	61.405 42.300 55.164 1.00 37.68 A	č
ATOM ATOM	5920 5921	CD LYS	760	61.620 41.455 53.916 1.00 40.82 A	C
ATOM	5922	CE LYS NZ LYS	760 760	62. 473 40. 229 54. 199 1. 00 42. 70 A	C
ATOM	5923	C LYS	760	62. 952 39. 600 52. 933 1. 00 44. 73 A 63. 885 44. 205 57. 396 1. 00 33. 20 A	N
ATOM	5924	0 LYS	760	63. 885 44. 205 57. 396 1. 00 33. 20 A 63. 874 43. 676 58. 504 1. 00 34. 38 A	C
ATOM	5925	N GLN	761	64. 914 44. 902 56. 939 1. 00 33. 26 A	O N
ATOM	5926	CA GLN	761	66. 106 45. 100 57. 744 1. 00 33. 22 A	C
ATOM ATOM	5927	CB GLN	761	67. 295 45. 422 56. 830 1. 00 35. 03 A	C
VION	5928	CG GLN	761	68. 638 45. 584 57. 525 1. 00 38. 28 A	С
			S	UBSTITUTE SHEET (RULE 26)	

					FΙ	G. 4	- 122	,		(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5929 5930 5931 5932 5933 5933 5934 5935 5938 5939 5940 5943 5944 5945 5947 5948 5949 5950 5951 5953 5956 5956	C O N CA C O CB SG N CA CB CCD1 CCE2 CZ C O N CA CB OG C O	GLN	761 761 761 761 762 762 762 762 762 763 763 763 763 763 763 763 763 764 764 764 764 764 764	68. 75 68. 48 69. 17 65. 81 66. 06 65. 27 64. 94 63. 88 63. 89 64. 47 63. 60 62. 993 61. 94 60. 618 59. 718 58. 80 59. 718 58. 139 62. 293 61. 499 63. 463 63. 907 65. 356 66. 215 63. 799 64. 195	9 46. 893 7 47. 969 7 46. 811 9 46. 251 4 46. 149 6 47. 337 5 48. 513 8 48. 216 2 48. 830 9 49. 643 6 51. 029 8 47. 271 8 46. 907 9 47. 892 9 49. 148 9 50. 300 9 48. 951 9 50. 202 9 45. 688 9 45. 276 9 45. 102 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 688 9 45. 276 9 45. 288 9 45. 288	58. 283 57. 739 59. 544 58. 701 59. 898 58. 161 58. 953 60. 023 61. 087 58. 025 58. 843 59. 742 60. 694 59. 981	1.00 42.12 1.00 43.62 1.00 44.19 1.00 32.55 1.00 32.49 1.00 32.03 1.00 33.26	A A A A A A A A A A A A A A A A A A A	Continued) C O N C O N C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM	5957 5958 5959 5960 5961	N CA CB CG	LEU LEU LEU LEU LEU	765 765 765 765 765 765	63. 264 63. 092 61. 624 61. 332 61. 996	43. 412 43. 716	64. 335 65. 747 66. 017 67. 299	1.00 48.04 1.00 51.59 1.00 50.97 1.00 50.79	A A A	N . C C C
ATOM ATOM ATOM ATOM ATOM	5962 5963 5964 5965 5966	CD2 C 0 N CD	LEU LEU LEU PRO PRO	765 765 765 766 766	59. 834 63. 533 62. 866 64. 667 65. 545	40. 213 44. 996 42. 588 41. 557 42. 776 43. 960	67. 221 67. 481 66. 676 66. 779 67. 372 67. 317	1. 00 50. 85 1. 00 50. 72 1. 00 54. 72 1. 00 55. 73 1. 00 57. 13 1. 00 57. 88	A A A A A	C C C O N C
ATOM ATOM ATOM ATOM ATOM ATOM	5971	CA CB CG C O OXT	PRO PRO PRO PRO PRO	766 766 766 766 766	65. 204 66. 600 66. 386 64. 352 63. 341	41. 775 42. 309 43. 797 41. 639 42. 370	68. 301 68. 604 68. 568 69. 565 69. 681	1.00 58.61 1.00 58.49 1.00 58.47 1.00 60.07 1.00 60.04	A A A A	C C C O
TER ATOM ATOM ATOM ATOM	5973 5974 5975 5976		PRO ASP ASP ASP	766 766 38 38 38 38	95. 909 96. 954 96. 905 97. 816	40. 805 45. 132 46. 047 47. 269 45. 544		1.00 61.88 1.00 32.66 1.00 32.61 1.00 30.88 1.00 31.65	A B B B	0 C C 0 0

			F I G. 4 - 1 2 3 (Cont	inued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	5979 O ASP 5980 N ASP 5981 CA ASP 5982 N SER 5983 CA SER 5984 CB SER 5985 OG SER 5986 C SER 5987 O SER 5988 N ARG 5989 CA ARG 5990 CB ARG 5991 CG ARG 5991 CG ARG 5991 CG ARG 5992 CD ARG 5993 NE ARG 5994 CZ ARG 5995 NH1 ARG 5996 NH2 ARG 5997 C ARG 5998 O ARG 5998 N LYS 6000 CA LYS 6001 CB LYS 6001 CB LYS 6001 CB LYS 6001 CB LYS 6002 CG LYS 6001 CB LYS 6002 CG LYS 6004 CE LYS 6005 NZ LYS 6006 C LYS 6007 O LYS 6006 C LYS 6007 O LYS 6007 O LYS 6008 N THR 6010 CB THR 6010 CB THR 6011 OG1 THR 6011 CG1 THR 6011 CG1 THR	38 38 39 39 39 39 39 40 40 40 40 40 40 41 41 41 41 41 41 42 42 42 42 42 42 42 42	94. 533	inued)
ATOM ATOM ATOM	6015 N TYR 6016 CA TYR 6017 CB TYR	43 43 43	94. 045 50. 750 67. 116 1. 00 20. 55 B N	
ATOM ATOM	6018 CG TYR 6019 CD1 TYR	43 43	95. 233 51. 153 65. 020 1. 00 20. 32 B C 95. 516 52. 062 63. 853 1. 00 19. 92 B C 94. 888 51. 863 62. 629 1. 00 22. 19 B C	
ATOM ATOM ATOM	6020 CE1 TYR 6021 CD2 TYR 6022 CE2 TYR	43 43	95. 133 52. 694 61. 546 1. 00 21. 23 B C 96. 403 53. 126 63. 970 1. 00 21. 09 B C	
ATOM ATOM	6023 CZ TYR 6024 OH TYR	43 43 43	90. 013	
ATOM ATOM	6025 C TYR 6026 O TYR	43 43	90. 247 54. 553 60. 600 1. 00 25. 44 B 0 92. 770 51. 631 65. 349 1. 00 18. 52 B C 92. 396 50. 640 64. 725 1. 00 17. 41 B 0	

			F I G. 4	- 124		(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6028 CA 6029 CB 6030 OG1 6031 CG2 6032 C 6033 O 6034 N 6035 CA 6036 CB 6037 CG 6038 CD1 6039 CD2 6040 C 6041 O 6042 N 6043 CA 6044 CB 6045 OG1 6046 CG2 6047 C 6048 O 6049 N 6050 CA 6051 CB 6052 CG 6053 OD1 6054 OD2 6055 C 6056 O 6057 N 6058 CA 6057 N 6058 CA 6059 CB 6060 CG 6061 CD1 6062 CE1 6062 CE1	LEU 45 LEU 45 THR 46 ASP 47 ASP 48 TYR 48 TYR 48 TYR 48 TYR 48	92. 007 52. 709 90. 633 52. 803 89. 762 53. 748 90. 195 55. 096 89. 875 53. 409 90. 521 53. 310 91. 511 53. 741 89. 296 53. 277 89. 026 53. 749 87. 570 53. 489 87. 163 54. 032 88. 050 53. 417 85. 698 53. 720 89. 300 55. 240 89. 827 55. 743 88. 948 55. 945 89. 156 57. 382 88. 550 57. 988 87. 148 57. 700 88. 745 59. 497 90. 634 57. 749 90. 999 58. 759 91. 491 56. 945 92. 910 57. 253 93. 731 56. 273 93. 365 56. 322 93. 116 57. 430 93. 339 55. 244 93. 357 57. 178 94. 057 58. 065 92. 951 56. 124 93. 332 55. 998 92. 823 54. 676 92. 867 54. 612 94. 062 54. 787 94. 098 54. 734	65. 532	B B B B B B B B B B B B B B B B B B B	
ATOM ATOM ATOM		TYR 48 TYR 48	91. 702 54. 383 91. 726 54. 329 92. 925 54. 503	56. 885	B B B	C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6066 OH T 6067 C T 6068 O T 6069 N L 6070 CA L 6071 CB L 6072 CG L 6073 CD1 L 6074 CD2 L	YR 48 YR 48 YR 48 EU 49 EU 49 EU 49 EU 49 EU 49 EU 49	92. 942 54. 434 92. 795 57. 170 93. 547 57. 853 91. 497 57. 416 90. 885 58. 485 89. 359 58. 437 88. 688 57. 157 87. 188 57. 305 89. 094 56. 889 91. 391 59. 886	53. 452 1. 00 18. 40 58. 899 1. 00 21. 85 58. 207 1. 00 21. 92 58. 996 1. 00 23. 08 58. 223 1. 00 26. 78 58. 381 1. 00 28. 14 57. 872 1. 00 28. 75 57. 980 1. 00 28. 04 56. 420 1. 00 28. 33	B B B B B B B	O C C C C C C C

			FIG. 4-126	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	M 6126 CB LEU M 6127 CG LEU M 6128 CD1 LEU M 6129 CD2 LEU M 6130 C LEU M 6131 O LEU M 6132 N LYS M 6133 CA LYS M 6134 CB LYS M 6135 CG LYS M 6136 CD LYS M 6137 CE LYS M 6138 NZ LYS M 6139 C LYS M 6139 C LYS M 6139 C LYS M 6140 O LYS M 6141 N LEU M 6142 CA LEU M 6145 CD1 LEU M 6145 CD1 LEU M 6146 CD2 LEU M 6147 C LEU M 6148 O LEU M 6149 N TYR	55555555555555555555555555555555555555	104. 959 58. 926 59. 515 1. 00 24. 45 B 105. 025 57. 911 58. 382 1. 00 22. 51 B 104. 335 56. 575 58. 631 1. 00 23. 77 B 104. 287 55. 792 57. 336 1. 00 23. 51 B 105. 083 55. 796 59. 703 1. 00 22. 83 B 105. 773 60. 161 59. 135 1. 00 24. 19 B 105. 428 60. 867 58. 187 1. 00 23. 47 B 106. 824 60. 456 59. 886 1. 00 23. 25 B 107. 631 61. 603 59. 532 1. 00 23. 25 B 107. 850 62. 922 61. 697 1. 00 29. 15 B 108. 868 63. 560 62. 638 1. 00 31. 22 B 109. 235 65. 233 64. 439 1. 00 32. 59 B 109. 235 65. 233 64. 439 1. 00 32. 59 B 109. 247 61. 945 58. 330 1. 00 23. 24 B 108. 818 61. 196 58. 330 1. 00 23. 24 B 109.	(Continued) C C C C C C C C C C C C C C C C C C
ATOM ATOM	6162 CA SER 6163 CB SER	59 59 59	112. 085 64. 604 52. 909 1. 00 21. 33 B 112. 245 64. 839 51. 479 1. 00 22. 11 B 110. 920 65. 275 50. 852 1. 00 21. 08 B	N C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6164 OG SER 6165 C SER 6166 O SER 6167 N LEU 6168 CA LEU 6169 CB LEU 6170 CG LEU 6171 CD1 LEU 6172 CD2 LEU 6173 C LEU	59 59 59 60 60 60 60 60	109.985 64.212 50.843 1.00 24.94 B 113.293 65.895 51.191 1.00 21.64 B 113.099 67.064 51.491 1.00 23.87 B 114.404 65.485 50.602 1.00 21.76 B 115.449 66.436 50.273 1.00 23.50 B 116.752 66.062 50.986 1.00 22.27 B 117.406 64.737 50.612 1.00 18.62 B 118.176 64.900 49.320 1.00 17.05 B 118.338 64.313 51.724 1.00 19.95 B 115.656 66.478 48.762 1.00 24.93 B	C O N C C C C C

				FIG. 4-127	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6209 6210 6211	N ARG CA ARG CB ARG CC TRP CC	60 61 61 61 61 61 61 61 62 62 62 62 62 62 62 62 63 63 63 64 64 64 64 64 64	115. 176 65. 604 48. 029 1. 00 23. 79 116. 375 67. 495 48. 302 1. 00 26. 02 116. 634 67. 659 46. 881 1. 00 27. 11 115. 693 68. 728 46. 329 1. 00 32. 13 115. 779 68. 979 44. 833 1. 00 38. 27 115. 002 70. 243 44. 495 1. 00 41. 78 114. 937 70. 506 43. 063 1. 00 46. 51 114. 298 71. 543 42. 525 1. 00 49. 47 113. 671 72. 420 43. 307 1. 00 48. 74 114. 266 71. 693 41. 205 1. 00 26. 01 118. 875 67. 186 46. 676 1. 00 26. 36 118. 877 67. 186 46. 095 1. 00 26. 36 118. 877 67. 186 46. 095 1. 00 24. 48 121. 024 66. 244 45. 355 1. 00 20. 04 121. 095 65. 145 46. 365 1. 00 18. 16 121. 954 65. 092 47. 508 1. 00 14. 54 121. 639 63. 272 47. 528 1. 00 15. 77 122. 956 65	B O N C C C C C C C C C C C C C C C C C C
ATOM ATOM	6212 6213	C SER O SER	64 64	125. 848 70. 724 45. 679 1. 00 27. 92 H 125. 399 68. 343 44. 255 1. 00 31. 08 H 124. 630 67. 488 44. 691 1. 00 31. 36 H	3 C
ATOM ATOM ATOM	6214 6215 6216	N ASP CA ASP CB ASP	65 65 65	126. 712 68. 176 44. 236 1. 00 31. 42 127. 306 66. 947 44. 728 1. 00 32. 55	B C
ATOM ATOM ATOM	6217 6218 6219	CG ASP OD1 ASP OD2 ASP	65 65 65	129. 158 65. 286 44. 302 1. 00 35. 12 128. 446 64. 261 44. 158 1. 00 33. 02	B C C
ATOM ATOM ATOM	6220 6221	C ASP O ASP N HIS	65 65 66	130. 331 65. 259 44. 728 1. 00 37. 02 B 127. 636 67. 045 46. 211 1. 00 32. 66 B 128. 076 66. 069 46. 818 1. 00 31. 78 B 127. 399 68. 217 46. 796 1. 00 33. 06 B	C C
		-	-	20.000 00.211 40.130 1.00 00.00 D	N ·

					FI	G. 4	128			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6223 6224 6225 6226 6227 6228 6229 6230 6231 6232 6233 6234 6235 6236 6237 6238 6241 6242 6243 6244 6245 6247 6255 6257 6258 6257 6258 6259 6262 6262 6262 6263 6262 6263 6262 6263 6262 6263 6262 6263 6262 6263 6	NDI CEE CON CAB CC CD I CON CAB CC CD I CC C	GLU GLU TYR TYR TYR TYR	66 66 66 66 66 66 67 67 67 67 67 67 68 68 68 68 68 69 69 69 69 69 69 70 70 70 70	F I 127. 704 128. 892 130. 032 131. 260 129. 959 131. 092 131. 897 126. 547 126. 602 125. 505 124. 379 124. 457 125. 601 125. 745 126. 408 125. 207 123. 015 122. 872 120. 634 120. 347 120. 333 120. 300 119. 657 119. 312 121. 391 121. 379 120. 333 120. 300 119. 657 119. 961 118. 497 117. 492 116. 729 117. 545 116. 656 118. 187 116. 508 115. 057 115. 799 114. 910 114. 396 113. 544	68. 440 69. 402 69. 076 68. 562 69. 238 68. 420 69. 001 69. 008 69. 479 70. 067	48. 203 48. 329 47. 416 47. 669 46. 047 45. 498 46. 459 49. 016 50. 245 49. 593 49. 593 48. 583 47. 482 49. 116 49. 592 51. 094 51. 914 53. 282 51. 685 53. 847 55. 191 49. 759 49. 759 49. 759 49. 759 50. 789 49. 693 48. 586 47. 693 48. 586 47. 693 48. 586 50. 260 51. 590 52. 482 54. 592 54. 592	1.00 32.64 1.00 35.63 1.00 39.09 1.00 40.29 1.00 41.80 1.00 42.37 1.00 42.11 1.00 31.01 1.00 30.92 1.00 30.05 1.00 28.07 1.00 27.21 1.00 29.99 1.00 33.25 1.00 34.83 1.00 27.52 1.00 27.10 1.00 26.72 1.00 25.74 1.00 25.74 1.00 22.93 1.00 22.93 1.00 22.93 1.00 22.75 1.00 23.47 1.00 23.47 1.00 23.47 1.00 23.81 1.00 26.50 1.00 26.50 1.00 26.72 1.00 27.89 1.00 27.89 1.00 28.86 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78 1.00 29.78	BBBBBBBBBBBBBBBBBBBBBBBBBBBBBBBBBBBBBB	Continued) CCCCCNCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC
ATOM ATOM ATOM ATOM ATOM	6267 6268 6269 6270 6271	CD2	TYR TYR TYR TYR TYR	70 70 70 70	114. 553 113. 701 113. 199 112. 346	69. 847 69. 141 67. 918 67. 221	55. 842 56. 686 56. 276 57. 103	1.00 28.33 1.00 28.03 1.00 28.21 1.00 30.20	B B B	C C C
V I O'M	04/1	U	111	70	114.056	71. 796	52. 983	1.00 34.45	В	C

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			FIG. 4-129	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6272 O TYR 6273 N LYS 6274 CA LYS 6275 CB LYS 6276 CG LYS 6277 CD LYS 6277 CD LYS 6278 CE LYS 6279 NZ LYS 6280 C LYS 6281 O LYS 6282 N GLN 6283 CA GLN 6284 CB GLN 6285 CG GLN 6285 CG GLN 6287 OE1 GLN 6288 NE2 GLN 6289 C GLN 6290 O GLN 6291 N GLU 6292 CA GLU 6292 CA GLU 6293 CB GLU 6294 CG GLU 6295 CD GLU 6296 OE1 GLU 6297 OE2 GLU 6298 C GLU 6298 C GLU 6299 O GLU 6299 O GLU 6299 CA ASN 6301 CA ASN 6302 CB ASN 6301 CA ASN 6302 CB ASN 6303 CG ASN 6304 OD1 ASN 6305 ND2 ASN 6306 C ASN 6307 O ASN 6307 O ASN 6308 N ASN 6309 CA ASN 6309 CA ASN 6301 CG ASN 6301 CG ASN 6301 CG ASN 6301 CA ASN 6301 CB ASN 6302 CB ASN 6303 CG ASN 6304 OD1 ASN 6305 ND2 ASN 6306 C ASN 6307 O ASN 6307 O ASN 6308 N ASN 6309 CA ASN 6309 CA ASN 6309 CA ASN 6301 CB ASN 6301 CB ASN 6301 CB ASN 6302 CB ASN 6303 CG ASN 6304 OD1 ASN 6305 ND2 ASN 6306 C ASN 6307 O ASN 6307 O ASN 6308 N ASN 6309 CA ASN 6309 CA ASN 6309 CA ASN 6309 CA ASN 6301 CB ASN 6301 CB ASN 6302 CB ASN 6303 CG ASN 6304 OD1 ASN 6305 ND2 ASN	73 73 73 73 73 73 74 74 74 74 74 74 75 75 75 75 75 76 11 76 11 76 11	114.425 72.914 53.336 1.00 33.43 1 112.787 71.410 53.002 1.00 39.59 1 111.714 72.284 53.461 1.00 44.28 1 110.408 71.904 52.763 1.00 45.57 1	3 C 3 N 5 C 6 C 6 C 7 C 8 C 8 C 9

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				· · · · · · · · · · · · · · · · · · ·	FIC	G. 4-	130			(Continued)
ATOM ~	6321		ILE	76	112. 341	77. 009	48. 967	1.00 42.01	В	Ċ
ATOM	6322	C	ILE	76	115. 243	76.589	53.043	1.00 39.85	В	C
ATOM	6323	0	ILE	76	115. 758	77. 701	53.150	1.00 41.15	В	0
ATOM	6324	N	LEU	77	115.862	75. 472	53.400	1.00 36.42	В	N
ATOM	6325	CA	LEU	77	117. 208	75. 498	53.941	1.00 34.22	В	C
ATOM	6326	CB	LEU	77	117. 227	74. 901	55. 351	1.00 34.28	В	C
ATOM	6327	CG	LEU	77	116. 155	75. 359	56.346	1.00 34.54	В	C
ATOM	6328		LEU	77	116. 435	74. 728	57. 701	1.00 33.23	В	C
ATOM	6329		LEU	77	116. 149	76.874	56.460	1.00 34.45	B	C
ATOM	6330	C	LEU	77	118. 121	74. 683	53. 036	1.00 32.91	В	C
ATOM	6331	0	LEU	77	117.657	73. 821	52. 289	1.00 32.49	В	0
ATOM	6332	N	VAL	78 70	119.417	74.967	53. 103	1.00 30.72	В	N
ATOM	6333	CA	VAL	78 70	120.409	74. 253	52.308	1.00 29.87	В	C
ATOM	6334	CB	VAL	78 79	121. 227	75. 227	51.431	1.00 30.20	В	C
ATOM	6335		VAL	78 78	122.327	74.480	50.691	1.00 29.01	В	C
ATOM	6336 6337	C	VAL	78 78	120.311	75.906	50.448	1.00 31.37 1.00 28.37	В	C
ATOM ATOM	6338	0	VAL VAL	78 78	121.346 121.781	73. 523 74. 087	53. 263 54. 261	1.00 28.38	В •В	C 0
ATOM	6339	N	PHE	79	121. 660	72. 272	52. 956	1.00 26.56	В	N N
ATOM	6340	CA	PHE	79 79	121.000	71.496	53.821	1.00 24.85	В	
ATOM	6341	CB	PHE	79 79	121.807	70. 247		1.00 24.65	В	C
ATOM	6342	CG	PHE	79	120.680	70. 531	55. 296	1.00 24.45	В	C C
ATOM	6343		PHE	79	119.499	71. 120	54. 857	1.00 22.02	В	C
ATOM	6344		PHE	79	120. 789	70. 168	56.636	1.00 20.13	В	C
ATOM	6345		PHE	79	118. 448	71. 338	55. 733	1.00 20.35	В	Č
ATOM	6346		PHE	79	119. 749	70. 382	57. 513	1.00 16.96	В	Č ·
ATOM	6347	CZ	PHE	79	118. 573	70.967	57.065	1.00 10.30	В	Č
ATOM	6348	C	PHE	79	123. 815	71.036	53. 151	1.00 24.95	В	Č .
ATOM	6349	ŏ	PHE	79	123.841	70. 729	51.960	1.00 24.30	В	0
ATOM	6350	Ň	ASN	80	124. 876	70. 992	53.948	1.00 23.66	В	N ·
ATOM	6351	CA	ASN	80	126. 174	70. 518	53. 517	1.00 23.32	B	Č
ATOM	6352	CB	ASN	80	127. 276	71.307	54. 220	1.00 22.91	B	č
ATOM	6353	CG	ASN	80	128.653	70.689	54. 032	1.00 22.91	B	č
ATOM	6354		ASN	80	128.916	69.567		1.00 23.26	B	Ö
ATOM	6355		ASN	80	129.542	71.421	53.364	1.00 21.99	В	Ň
ATOM	6356	C	ASN	80	126.156	69.077	54.018	1.00 24.17	В	Ċ
ATOM	6357	0	ASN	80	126.168	68.842	55. 222	1.00 25.80	В	0
ATOM	6358	N	ALA	81	126.116	68.116	53.105	1.00 23.17	В	N
ATOM	6359	CA	ALA	81	126.054	66.713	53.496	1.00 24.07	В	C
ATOM	6360	CB	ALA	81	126.025	65.819	52.246	1.00 20.69	В	C
ATOM	6361	C	ALA	81	127.167	66.256	54.434	1.00 25.23	В	C
ATOM	6362	0	ALA	81	126.925	65.462	55.347	1.00 25.26	В	0
ATOM	6363	N	GLU	82	128.377	66.764	54. 222	1.00 26.73	В	N
ATOM	6364	CA	GLU	82	129.525	66.351	55.024	1.00 29.51	В	C
ATOM	6365	CB	GLU	82	130.820	66.835	54.361	1.00 32.02	В	C
ATOM	6366	CG	GLU	82	132.124	66.326	55.005	1.00 35.72	В	. C
ATOM	6367	CD	GLU	82	132. 287	64.800	54.955	1.00 38.90	В	C
ATOM	6368		GLU	82	132.064	64. 191	53.884	1.00 38.71	В	0
ATOM	6369	0E2	GLU	82	132.659	64. 209	55. 995	1.00 40.81	В	0

					FΙ	G. 4	- 1 3 1	L		(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6370 6371 6372 6373 6374 6375 6376 6376 6377 6378 6380 6381 6382 6383 6384 6389 6390 6391 6392 6393 6394 6403 6404 6405 6407 6407 6408 6407 6411 6412 6413 6416 6417 6418	O N CACE CCC CC CC O N CACE CO N CACE CCC CC	TYR TYR TYR TYR TYR TYR TYR	82 82 83 83 83 83 83 83 83 83 83 83 83 84 84 85 85 85 85 86 86 86 87 87 87 87 88 88 88 88 88 89 89 89 89 89 89 89 89	F I 129. 528 130. 102 128. 888 128. 877 129. 504 130. 821 131. 914 133. 120 130. 966 132. 162 133. 234 134. 413 127. 490 127. 340 126. 478 125. 136 124. 668 123. 511 125. 565 125. 201 126. 446 127. 356 128. 051 127. 338 124. 381 124. 432 123. 622 122. 787 121. 392 120. 734 122. 658 123. 307 121. 806 121. 530 122. 588 123. 307 121. 806 121. 530 122. 588 123. 387 120. 191 119. 832 119. 444 117. 357 116. 094 117. 006 118. 422 119. 235 117. 745 117. 901 118. 060	66. 757 66. 051 67. 872 68. 329 69. 722 69. 834 69. 049 70. 704 70. 791 70. 091 68. 355 68. 093 68. 684 68. 726	7 56. 497 57. 324 56. 834 58. 223 58. 320 57. 963 57. 971 56. 517 55. 815 56. 195 55. 486 58. 853 60. 044 58. 063 58. 601	7 1.00 29.17 4 1.00 28.55 4 1.00 29.07 8 1.00 28.95 9 1.00 30.17 1.00 33.40 1.00 33.79 1.00 36.07 1.00 36.91 1.00 38.12 1.00 42.42 1.00 28.16 1.00 29.04 1.00 25.68 1.00 24.77 1.00 24.95	B B B B B B B B B B B B B B B B B B B	CONCCCCCCCCONCCCONCCCONCCCONCCCCCONCCCCCONCCCC
				SL	JBSTITUTE				-	•

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			FIG. 4	- 1 3 3		(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6473 C 6474 O 6475 N 6476 CA 6477 CB 6478 CG 6479 OD1 6480 OD2 6481 C 6482 O 6483 N 6484 CA 6485 CB 6486 CG 6487 CD 6490 C 6491 O 6492 N 6493 CA 6494 CB 6495 CG 6496 CD1 H 6497 CD2 H 6497 CD2 H 6497 CD2 H 6497 CD2 H 6498 CE1 H 6499 CE2 H 6499 CE2 H 6490 CZ 6501 C 6502 O 6503 N	PHE 95 PHE 96 ASP 97 GLU 97 GL	107. 978 83. 33 107. 476 85. 29 109. 095 83. 09 108. 594 85. 06 109. 403 83. 96 104. 825 86. 10 103. 740 85. 78 104. 941 86. 68 103. 775 86. 96 104. 167 87. 78 104. 793 86. 94 104. 234 85. 87 105. 835 87. 36 102. 674 87. 71 101. 498 87. 40 103. 050 88. 70 102. 068 89. 49 102. 389 90. 99 102. 389 90. 99 102. 389 90. 99 102. 397 91. 55 103. 629 91. 14 103. 714 91. 49 104. 514 90. 46 101. 970 89. 12 101. 652 89. 97 102. 234 87. 85 102. 181 87. 39 102. 792 85. 43 102. 792 85. 43 103. 564 86. 07 102. 064 84. 30 103. 609 85. 59 102. 103 83. 82 102. 876 84. 46 100. 764 87. 448 100. 578 87. 544 99. 770 87. 383	26 48. 268 1. 00 51. 89 30 47. 005 1. 00 50. 37 31 46. 205 1. 00 50. 62 30 46. 441 1. 00 50. 62 30 46. 441 1. 00 55. 66 31 50. 639 1. 00 55. 66 32 50. 69 1. 00 55. 16 31 51. 835 1. 00 56. 69 4 52. 668 1. 00 57. 24 5 53. 900 1. 00 60. 82 6 55. 321 1. 00 60. 82 6 55. 538 1. 00 57. 24 1 52. 100 1. 00 58. 26 3 51. 130 1. 00 57. 07 6 50. 395 1. 00 57. 07 6 50. 395 1. 00 59. 15 3 51. 935 1. 00 61. 76 0 52. 729 1. 00 63. 57 0 52. 729 1. 00 63. 57 0 53. 927 1. 00 64. 73 3 48. 917 1. 00 56. 86 48. 598 1. 00 54. 75 44. 749 1. 00 52. 58 47. 117 1. 00 52. 58	B B B B B B B B B B B B B B B B B B B	(Continued) C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM	6505 C G 6506 O G	GLY 99 GLY 99 GLY 99	98. 383 87. 441 97. 918 86. 192 97. 020 86. 246	46. 376 1. 00 47. 41 45. 540 1. 00 48. 42	B B B	C C 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6508 CA H 6509 CB H 6510 CG H 6511 CD2 H 6512 ND1 H 6513 CE1 H 6514 NE2 H 6515 C H	IIS 100 IIS 100 IIS 100 IIS 100	98. 530 85. 065 98. 200 83. 780 98. 787 83. 686 98. 004 84. 414 98. 345 85. 437 96. 711 84. 075 96. 288 84. 857 97. 262 85. 691 98. 822 82. 677	46. 104 1. 00 43. 24 44. 694 1. 00 41. 93 43. 651 1. 00 39. 37 42. 833 1. 00 38. 83 43. 321 1. 00 39. 65 42. 344 1. 00 38. 90 42. 029 1. 00 38. 71 46. 940 1. 00 42. 56	B B B B B	N C C C C N C N C
UIOII	6516 O H	IS 100	99.916 82.846	47. 473 1. 00 43. 12	В	0

	FIG. 4-134	(Continued)
ATOM 6518 CA SER ATOM 6518 CA SER ATOM 6519 CB SER ATOM 6520 OG SER ATOM 6521 C SER ATOM 6522 O SER ATOM 6523 N ILE ATOM 6524 CA ILE ATOM 6525 CB ILE ATOM 6526 CG2 ILE ATOM 6527 CG1 ILE ATOM 6528 CD1 ILE ATOM 6529 C ILE ATOM 6530 O ILE ATOM 6531 N ASN ATOM 6531 N ASN ATOM 6532 CA ASN ATOM 6533 CB ASN ATOM 6534 CG ASN ATOM 6535 OD1 ASN ATOM 6536 ND2 ASN ATOM 6537 C ASN ATOM 6538 O ASN ATOM 6538 O ASN ATOM 6539 N ASP ATOM 6539 N ASP ATOM 6540 CA ASP ATOM 6541 CB ASP ATOM 6542 CG ASP ATOM 6543 OD1 ASP ATOM 6544 OD2 ASP ATOM 6545 C ASP ATOM 6545 C ASP ATOM 6546 O ASP ATOM 6547 N TYR ATOM 6548 CA TYR ATOM 6549 CB TYR ATOM 6550 CG TYR ATOM 6551 CD1 TYR ATOM 6551 CD1 TYR ATOM 6555 CZ TYR ATOM 6556 OH TYR ATOM 6557 C TYR ATOM 6557 C TYR ATOM 6557 C TYR ATOM 6559 N SER ATOM 6559 N SER ATOM 6559 N SER	FIG. 4 - 134 101 98.139 81.547 47.063 1.00 41.90 101 98.716 80.442 47.817 1.00 43.20 101 97.623 79.527 48.382 1.00 43.41 101 96.852 78.931 47.354 1.00 44.00 101 99.582 79.680 46.820 1.00 42.92 101 99.083 79.213 45.794 1.00 41.90 102 100.880 79.584 47.095 1.00 41.90 102 101.762 78.874 46.183 1.00 42.10 102 103.255 79.286 46.369 1.00 43.10 102 103.370 80.811 46.404 1.00 43.52 102 103.824 78.700 47.660 1.00 45.01 102 105.294 79.038 47.895 1.00 46.96 102 101.598 77.380 46.415 1.00 41.08 102 101.677 76.901 47.544 1.00 41.27 103 101.342 76.648 45.339 1.00 40.05 103 101.157 75.211 45.434 1.00 39.20 103 100.502 74.674 44.163 1.00 39.98 103 100.190 73.199 44.257 1.00 39.82 103 100.866 72.396 43.448 1.00 40.83 103 100.866 72.396 43.448 1.00 40.75 103 102.486 74.508 45.645 1.00 37.42 104 104.808 74.303 44.982 1.00 34.14 104 104.819 72.955 44.248 1.00 33.77 104 104.808 74.303 44.982 1.00 34.77 104 106.635 72.272 44.655 1.00 34.77 104 106.835 72.525 45.453 1.00 33.54 105 107.103 74.985 44.607 1.00 33.77 104 106.861 70.919 44.178 1.00 33.77 104 106.061 70.919 44.178 1.00 33.77 104 105.827 75.253 44.607 1.00 33.54 105 107.103 74.985 44.607 1.00 33.77 104 105.867 72.072 44.655 1.00 33.77 104 105.867 75.254 44.082 1.00 33.75 105 108.867 75.824 44.082 1.00 33.54 105 107.103 74.985 44.607 1.00 32.32 105 108.167 75.824 44.082 1.00 33.54 105 107.103 74.985 44.607 1.00 32.32 105 108.167 75.824 44.082 1.00 33.54 105 107.103 74.985 44.607 1.00 32.32 105 108.867 75.824 44.082 1.00 33.54 105 109.515 75.662 46.218 1.00 35.82 105 109.387 74.282 48.208 1.00 36.01 105 107.103 74.985 44.009 1.00 36.01 105 107.103 74.985 44.009 1.00 36.01 105 109.387 74.282 48.208 1.00 38.47 105 109.387 74.282 48.208 1.00 38.47 105 109.387 74.282 48.208 1.00 38.47 105 109.048 73.754 48.065 1.00 37.17 105 109.048 73.754 48.065 1.00 37.17 105 109.048 73.754 48.065 1.00 38.47	(Continued) B N B C B C B C B C B C B C B C B C B C B C
ATOM 6561 CB SER ATOM 6562 OG SER ATOM 6563 C SER ATOM 6564 O SER ATOM 6565 N ILE	106 110.648 74.391 40.758 1.00 24.49 106 111.662 74.145 39.806 1.00 24.16 106 112.341 75.926 41.745 1.00 26.32 106 112.168 76.821 40.919 1.00 28.04	B C B O B C

	FIG. 4-135	(Continued)
ATOM 6594 N GLY ATOM 6595 CA GLY ATOM 6596 C GLY ATOM 6597 O GLY ATOM 6598 N GLN ATOM 6599 CA GLN ATOM 6699 CA GLN ATOM 6600 CB GLN ATOM 6601 CG GLN ATOM 6602 CD GLN ATOM 6603 OE1 GLN ATOM 6604 NE2 GLN ATOM 6605 C GLN ATOM 6606 O GLN ATOM 6606 O GLN ATOM 6607 N PHE ATOM 6608 CA PHE ATOM 6609 CB PHE ATOM 6610 CG PHE ATOM 6611 CD1 PHE ATOM 6612 CD2 PHE ATOM 6613 CE1 PHE	107	$\tt CCCCCCONCCOCONCCCCCONCCCONCCCONCCCONCC$

				FIG. 4-136	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6650 6651 6652 6653 6654 6655	C PHE O PHE N ILE CA ILE CB ILE CG2 ILE CG1 ILE CG2 ILE CG3 ILE CG3 ILE CG3 ILE CG3 ILE CG4 ILE CG5 ILE CG6 ILE CG7 ILE CG7 ILE CG7 ILE CG8 ILE CG9 IL	113 114 114 114 114 114 115 115 115 115 115	117. 386 85. 152 36. 819 1.00 35. 71 B 114. 831 80. 896 41. 058 1.00 30. 65 B 115. 308 79. 829 41. 425 1.00 30. 90 B 113. 557 81. 205 41. 219 1.00 30. 99 B 112. 630 80. 258 41. 791 1.00 29. 81 B 112. 394 80. 504 43. 293 1.00 28. 60 B 111. 911 81. 915 43. 529 1.00 30. 57 B 111. 336 79. 367 45. 325 1.00 30. 57 B 111. 336 80. 403 41. 019 1.00 29. 79 B 110. 895 81. 508 40. 715 1. 00 28. 83 B 110. 756 79. 265 40. 671 1. 00 30. 43 B 109. 516 79. 223 39. 925 1. 00 29. 05 B 109. 596 78. 108 38. 890 1. 00 28. 22 B 108. 449 77. 898 37. 912 1. 00 28. 22 B 108. 427 79. 001 36. 872 1. 00 29. 59 B 108. 427	CCONCCCCONCCCCCONCCCCONCCCCONCCCCONCCCC
ATOM ATOM ATOM	6655 6656 6657	CD1 TYR CE1 TYR CD2 TYR	118 118	101. 256 81. 301 41. 092 1. 00 30. 04 B 101. 355 82. 373 40. 210 1. 00 31. 36 B	C ·
ATOM ATOM	6658 6659	CE2 TYR CZ TYR	118 118 118	98. 915 81. 703 41. 316 1. 00 30. 41 B 99. 003 82. 768 40. 439 1. 00 31. 17 B 100. 222 83. 101 39. 891 1. 00 31. 56 B	C C C
ATOM ATOM ATOM	6660 6661 6662	OH TYR C TYR O TYR	118 118 118	100. 298 84. 179 39. 039 1. 00 33. 43 B 98. 814 78. 240 41. 038 1. 00 27. 66 B	0 C
ATOM		N ASN	119	99. 046 77. 917 39. 874 1. 00 26. 73 B 97. 582 78. 450 41. 499 1. 00 27. 22 B	O N

					FIG. 4-137	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6664 6665 6666 6667 6668 6670 6673 6674 6675 6676 6680 6681 6682 6683 6684 6685 6688 6689 6691 6692 6693 6699 6699 6690 6701	CBCCODNDCONCABCCCCCONCACCCCCCCCCCCCCCCCCCCCCCCCC	ASN ASN ASN ASN TYR	119 119 119 119 120 120 120 120 120 120 120 121 121 121	96. 397 78. 261 40. 659 1. 00 27. 10 86. 422 79. 203 39. 449 1. 00 27. 22 895. 918 80. 599 39. 777 1. 00 27. 62 894. 905 80. 761 40. 456 1. 00 26. 76 80. 6613 81. 612 39. 277 1. 00 25. 87 896. 342 76. 810 40. 171 1. 00 27. 88 895. 923 76. 534 39. 045 1. 00 27. 93 896. 771 75. 888 41. 028 1. 00 27. 57 86. 795 74. 466 40. 702 1. 00 29. 01 897. 396 73. 669 41. 866 1. 00 30. 85 897. 421 72. 171 41. 635 1. 00 32. 83 898. 466 71. 568 40. 940 1. 00 33. 76 88. 484 70. 190 40. 717 1. 00 35. 03 85 96. 389 71. 358 42. 100 1. 00 34. 41 896. 394 69. 981 41. 880 1. 00 34. 35 897. 444 69. 403 41. 191 1. 00 35. 47 897. 462 68. 039 40. 987 1. 00 35. 56 895. 431 73. 863 40. 364 1. 00 29. 17 894. 458 74. 034 41. 099 1. 00 31. 09 895. 368 73. 148 39. 248 1. 00 27. 53 894. 136 72. 487 38. 842 1. 00 25. 45 894. 136 72. 487 38. 842 1. 00 25. 41 8 894. 124 70. 082 38. 977 1. 00 24. 18 894. 124 70. 082 38. 977 1. 00 24. 18 894. 124 70. 082 38. 977 1. 00 24. 18 894. 124 70. 082 38. 977 1. 00 24. 18 894. 124 70. 082 38. 977 1. 00 24. 18 894. 124 70. 082 38. 977 1. 00 24. 18 894. 124 70. 188 894. 124 70. 18	(Continued) C C C C C C C C C C C C C C C C C C
ATOM ATOM	6702 6703	CG CD	GLN GLN	123 123	93. 689 66. 433 32. 948 1. 00 23. 42 B 94. 242 66. 857 31. 591 1. 00 24. 37 B	C C
ATOM ATOM ATOM	6704 6705 6706	NE2 C	GLN GLN GLN	123 123 123	95. 399 66. 606 31. 275 1. 00 26. 71 B 93. 402 67. 493 30. 779 1. 00 23. 80 B 93. 856 65. 194 35. 805 1. 00 20. 06 B	O N C
ATOM ATOM ATOM	6707 6708 6709	O N CA	GLN TRP TRP	123 124 124	93. 258 64. 741 36. 786 1. 00 17. 04 B 94. 630 64. 438 35. 030 1. 00 17. 49 B	0 N .
ATOM ATOM	6710 6711	CB CG	TRP TRP	124 124	95. 165 62. 298 33. 984 1. 00 16. 19 B 94. 351 62. 735 32. 797 1. 00 18. 11 B	C C C
ATOM	6712	CD2	IKP	124	92. 939 63. 014 32. 764 1. 00 17. 55 B	Č

	•			•						(Continued)
					FIG	G. 4-	138	•		(Continued)
. = 0										
ATOM	6713		TRP	124	92.630	63.449	31.455	1.00 16.84	В	C
ATOM	6714	CE3		124	91.909	62.942	33.713	1.00 17.02	В	C
ATOM	6715	CD1		124	94. 819	62.999	31.539	1.00 19.00	В	C
ATOM	6716	NE1		124	93. 794	63.429	30.731	1.00 18.26	В	N
ATOM	6717		TRP	124	91.331	63.815	31.067	1.00 15.16	В	C
ATOM	6718		TRP	124	90.615	63. 305	33. 326	1.00 16.85	В	C .
ATOM	6719		TRP	124	90. 342	63. 737	32.011	1.00 16.12	В	C
ATOM	6720	C	TRP	124	95. 718	62.679	36.427	1.00 17.28	В	C
ATOM	6721	0	TRP	124	95.816	63. 437	37. 397	1.00 17.74	В	0
ATOM	6722	N	ARG	125	96. 430	61.560	36. 339	1.00 15.31	В	N
ATOM	6723	CA	ARG	125	97.317	61. 185	37. 429	1.00 16.66	В	C
ATOM	6724	CB	ARG	125	97.666	59. 702	37. 323	1.00 16.96	В	C
ATOM	6725	CG	ARG	125	98. 908	59. 288	38.076	1.00 18.35	В	C
ATOM	6726	CD	ARG	125			38.794	1.00 18.85	В	C
ATOM	6727	NE CZ	ARG	125	98. 049	56. 965	37.972	1.00 18.57	В	N
ATOM	6728	CZ	ARG	125	97.547	55. 842	38. 475	1.00 17.58	В	C
ATOM	6729			125	96.972	54. 944	37.693	1.00 16.96	В	N
ATOM ATOM	6730 6731	C	ARG	125	97.626	55.621	39.776	1.00 17.03	B	N
ATOM	6732	0	ARG ARG	125	98.582	62.027	37. 568	1.00 18.54	В	C
ATOM	6733	N	HIS	125 126	99. 075 99. 099	62. 227	38.674	1.00 18.06	В	0
ATOM	6734	CA	HIS	126	100. 300	62. 533 63. 353	36.454	1.00 20.06	В	N
ATOM	6735	CB	HIS	126	100.300	62.673	36. 487 35. 673	1.00 18.20 1.00 18.72	B	C
ATOM	6736	CG	HIS	126	101.721	61. 295	36. 151	1.00 18.72	B B	C
ATOM	6737		HIS	126	101.519	60.084	35. 581	1.00 19.88	В	C
ATOM	6738		HIS	126	102. 341	61.054	37. 360	1.00 20.00	В	· N
ATOM	6739		HIS	126	102.510	59. 753	37. 512	1.00 19.55	В	C
ATOM	6740		HIS	126	102.019	59. 142	36. 447	1.00 22.65	В	N
ATOM	6741	С	HIS	126	100.079	64. 772	35. 966	1.00 18.28	В	Č
ATOM	6742	0	HIS	126	100.692	65.716	36. 462	1.00 18.27	B	ŏ
ATOM	6743	N	SER	127	99. 204	64.921	34.974	1.00 16.08	B	Ň
ATOM	6744	CA	SER	127	98.936	66. 230	34. 382	1.00 16.78	B	Ċ
ATOM	6745	CB	SER	127	98. 209	66.070	33.037	1.00 15.96	B	č
ATOM	6746	0G	SER	127	96.999	65. 349	33.179	1.00 17.80	В	Ō
ATOM	6747	C	SER	127	98. 151	67. 203	35. 261	1.00 16.75	В	Č
ATOM	6748	0	SER	127	97. 523	66. 81 6	36. 247	1.00 17.88	В	0
ATOM	6749	N	TYR	128	98. 205	68.473	34.873	1.00 15.65	В	N
ATOM	6750	CA	TYR	128	97.520	69.556	35. 559	1.00 17.91	В	С
ATOM	6751	CB	TYR	128	97.815	69.506	37.060	1.00 17.70	В	С
ATOM	6752	CG	TYR	128	99.253	69. 796	37. 444	1.00 17.20	В	С
ATOM	6753	CD1	TYR	128	99. 725	71. 107	37. 540	1.00 16.17	В	С
ATOM	6754		TYR	128	101.036	71.375	37. 927	1.00 16.04	В	С
ATOM	6755		TYR	128	100. 135	68.759	37. 739	1.00 17.12	В	C .
ATOM	6756		TYR	128	101.449	69.016	38. 123	1.00 15.90	В	C
ATOM	6757	CZ	TYR	128	101.891	70.322	38. 216	1.00 17.19	В	С
ATOM	6758	OH C	TYR	128	103. 190	70.572	38. 603	1.00 20.16	В	0
ATOM	6759	C	TYR	128	97. 977	70.897	34. 992	1.00 19.77	В	C
ATOM	6760	0 N	TYR	128	98. 970	70.972	34. 268	1.00 21.70	В	0
ATOM	6761	N	THR	129	97. 239	71.955	35. 291	1.00 20.48	В	N

	·								
				FI	G. 4	- 139)		(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6763 6764 6765 6766 6767 6778 6779 6778 6778 6778 6778	ALA SER SER SER SER SER SER SER TYR SER TYR TYR TYR TYR TYR TYR TYR ASP	129 129 129 129 130 130 130 130 131 131 131 131 131 132 132 132 132 132	97. 647 96. 599 95. 353 96. 428 97. 856 97. 462 98. 754 99. 789 99. 269 99. 514 99. 934 99. 056 97. 713 101. 290 101. 448 102. 272 103. 611 104. 558 104. 179 103. 082 102. 696 104. 887 104. 510 103. 415 103. 023 104. 143 105. 674 104. 954 104. 954 103. 732 104. 954 103. 732 107. 385 108. 031 109. 444 110. 267	73. 276 73. 968 74. 045 73. 213 74. 136 73. 765 75. 289 76. 222 75. 631 77. 525 77. 632 78. 523 79. 818 80. 775 79. 851 79. 334 80. 506 79. 634 77. 721 76. 383 77. 250 75. 911 76. 383 77. 250 75. 911 75. 486 74. 171 81. 929 82. 790 82. 165 83. 465 84. 287 85. 008 84. 332 86. 253 83. 228 84. 382 86. 253 83. 764 83. 750	34. 840 33. 950 34. 652 32. 634 36. 069 37. 182 35. 854 36. 926 37. 859 36. 796 37. 333 36. 796 37. 347 36. 519 36. 519 36. 516 35. 834 37. 283 37. 383 36. 913 37. 463 37. 383 36. 792 37. 347 36. 519 36. 636 37. 383 37. 199 36. 516 37. 383 37. 283 37. 383 37. 283 37. 383 37. 283 37. 383 37. 283 37. 283 37. 383 37. 283 37. 383 37. 38	1.00 22.26 1.00 23.04 1.00 24.93 1.00 22.70 1.00 22.23 1.00 20.98 1.00 22.77 1.00 23.41 1.00 19.73 1.00 26.66 1.00 27.20 1.00 30.14 1.00 30.56 1.00 32.67 1.00 31.00 1.00 32.67 1.00 31.40 1.00 26.31 1.00 26.45 1.00 26.31 1.00 26.45 1.00 26.58 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.63 1.00 26.74	B B B B B B B B B B B B B B B B B B B	$\begin{array}{l} \texttt{C} \\ \texttt{C} \\ \texttt{O} \\ \texttt{C} \\ \texttt{C} \\ \texttt{O} \\ \texttt{C} \\ $
ATOM ATOM	6802 CG 6803 CG	2 ILE 1 ILE	134 134	111.718 109.649	83. 392 82. 737	37. 593 36. 312	1.00 31.90 1.00 32.72	B B	
ATOM ATOM ATOM	6805 C 6806 O	ILE ILE ILE	134 134 134	109. 887 109. 521	82. 794 84. 911 86. 065	34. 909 39. 483 39. 261	1.00 31.29 1.00 34.02 1.00 33.25	B B B	C C
ATOM ATOM ATOM	6807 N 6808 CA 6809 CB	TYR	135 135 135	111.167 110.657		40. 507 41. 475 42. 868	1.00 35.09 1.00 36.09 1.00 36.02	B B B	N C C C
ATOM	6810 CG	TYR	135	111. 222	86.011	44.000	1.00 36.66	В	C

										(Continued)
					FIC	3.4-	140			
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6811 6812 6813 6814 6815 6816	CE1 CD2 CE2 CZ OH C	TYR TYR TYR TYR TYR TYR TYR	135 135 135 135 135 135	110. 635 111. 134 112. 332 112. 839 112. 235 112. 740 112. 688	87. 222 87. 971 85. 573 86. 316 87. 515 88. 258 85. 511	44. 363 45. 424 44. 729 45. 786 46. 131 47. 179 41. 470	1.00 34.73 1.00 34.55 1.00 35.12 1.00 35.07 1.00 35.31 1.00 35.05 1.00 38.19	B B B B B	C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6818 6819 6820 6821 6822 6823 6824 6825		TYR ASP ASP ASP ASP ASP ASP	135 136 136 136 136 136 136	113. 293 113. 304 114. 759 115. 187 116. 690 117. 107 117. 456 115. 316	84. 517 86. 600 86. 692 87. 969 88. 051 88. 577 87. 602 86. 679	41. 873 41. 014 40. 965 40. 237 40. 030 38. 978 40. 911 42. 382	1.00 37.81 1.00 40.56 1.00 42.09 1.00 42.45 1.00 43.61 1.00 45.53 1.00 41.77 1.00 43.14	B B B B B	0 N C C C 0 0
ATOM ATOM ATOM ATOM ATOM ATOM	6826 6827 6828 6829 6830 6831 6832	CD2	ASP LEU LEU LEU LEU LEU	136 137 137 137 137 137	114. 972 116. 181 116. 761 117. 219 116. 058 116. 582 115. 199	87. 522 85. 713 85. 577 84. 135 83. 136 81. 716 83. 291	43. 209 42. 656 43. 978 44. 182 44. 117 43. 991 45. 361	1.00 42.49 1.00 44.92 1.00 48.26 1.00 48.88 1.00 49.07 1.00 50.17 1.00 48.91	B B B B B	O N C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6833 6834 6835 6836 6837 6838 6839 6840	C O N CA CB CG OD1 ND2	LEU ASN ASN ASN ASN ASN	137 137 138 138 138 138 138	117. 908 118. 309 118. 429 119. 522 120. 330 120. 728 121. 232 120. 512	86. 544 86. 750 87. 139 88. 096 88. 151 86. 775 85. 945 86. 530	44. 228 45. 370 43. 160 43. 280 41. 983 41. 484 42. 244 40. 194	1.00 50.19 1.00 51.45 1.00 52.26 1.00 53.21 1.00 54.36 1.00 56.39 1.00 57.23 1.00 56.67	B B B B B	C O N C C C O N
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6841 6842 6843 6844 6845 6846 6847	C O N CA CB CG CD	ASN ASN LYS LYS LYS LYS LYS	138 138 139 139 139 139	118. 935 119. 259 118. 064 117. 417 116. 807 117. 726 116. 996	89. 472 90. 101 89. 929 91. 228 91. 657 91. 520 91. 874	43. 567 44. 571 42. 675 42. 814 41. 480 40. 290 39. 006	1. 00 54. 11 1. 00 54. 39 1. 00 55. 06 1. 00 56. 16 1. 00 56. 75 1. 00 58. 34 1. 00 59. 63	B B B B B	C O N C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6848 6849 6850 6851 6852 6853 6854 6855	CE NZ C O N CA CB CG	LYS LYS LYS LYS ARG ARG ARG ARG	139 139 139 139 140 140 140	117. 887 117. 196 116. 302 115. 669 116. 061 114. 994 115. 433 116. 063	91. 650 91. 995 91. 183 92. 202 90. 006 89. 838 90. 341 89. 260	37. 793 36. 518 43. 857 44. 139 44. 425 45. 409 46. 787 47. 649	1. 00 61. 32 1. 00 62. 59 1. 00 56. 78 1. 00 57. 22 1. 00 57. 14 1. 00 57. 44 1. 00 58. 40 1. 00 61. 65	B B B B B	C N C O N C C
ATOM ATOM ATOM ATOM	6856 6857 6858 6859	CD NE CZ NH1	ARG ARG ARG	140 140 140 140	116. 091 116. 578 115. 979 114. 857	89. 658 88. 575 87. 394 87. 124	49. 116 49. 972 50. 112 49. 453	1. 00 64. 17 1. 00 67. 20 1. 00 68. 02 1. 00 68. 21	B B B B	C C N C N

	FIG. 4-141	(Continued)
ATOM 6861 C ARG 1 ATOM 6862 O ARG 1 ATOM 6863 N GLN 1 ATOM 6864 CA GLN 1 ATOM 6865 CB GLN 1 ATOM 6866 CG GLN 1 ATOM 6867 CD GLN 1 ATOM 6868 OE1 GLN 1 ATOM 6869 NE2 GLN 1 ATOM 6870 C GLN 1 ATOM 6871 O GLN 1	108. 196	B

				FIG. 4-142	(Continued)
ATOM	6909		146	107. 685 88. 081 26. 543 1. 00 45: 03 B	0
ATOM	6910		146	107.641 90.269 26.387 1.00 45.44 B	
ATOM	6911	C GLU	146	106. 978 87. 241 30. 821 1. 00 46. 25 B	
ATOM ATOM	6912 6913	0 GLU N ARG	146	107. 805 86. 334 30. 912 1. 00 47. 62 B	
ATOM	6914	CA ARG	147 147	105. 823 87. 221 31. 474 1. 00 44. 79 B 105. 475 86. 119 32. 360 1. 00 43. 34 B	
ATOM	6915	CB ARG	147	105. 475 86. 119 32. 360 1. 00 43. 34 B 104. 469 86. 595 33. 410 1. 00 44. 21 B	
ATOM	6916	CG ARG	147	104. 998 87. 678 34. 320 1. 00 46. 85 B	C C
ATOM	6917	CD ARG	147	103. 995 88. 007 35. 410 1. 00 49. 84 B	Č
ATOM	6918	NE ARG	147	102. 805 88. 651 34. 866 1. 00 53. 22 B	Ň
ATOM	6919	CZ ARG	147	101.733 88.970 35.584 1.00 54.21 B	C
ATOM	6920	NH1 ARG	147	101. 696 88. 699 36. 884 1. 00 53. 97 B	N
ATOM ATOM	6921 6922	NH2 ARG	147	100. 701 89. 569 34. 999 1. 00 54. 56 B	N
ATOM	6923	C ARG O ARG	147 147	104. 905 84. 894 31. 648 1. 00 41. 06 B 104. 304 84. 996 30. 580 1. 00 41. 00 B	C
ATOM	6924	N ILE	148	104. 304 84. 996 30. 580 1. 00 41. 00 B 105. 103 83. 732 32. 259 1. 00 38. 31 B	0
ATOM	6925	CA ILE	148	104. 590 82. 485 31. 721 1. 00 35. 74 B	N C
ATOM	6926	CB ILE	148	105. 019 81. 305 32. 616 1. 00 35. 07 B	Č
ATOM	6927	CG2 ILE	148	104. 458 79. 996 32. 073 1. 00 34. 22 B	č
ATOM	6928	CG1 ILE	148	106. 549 81. 255 32. 679 1. 00 33. 62 B	C
ATOM ATOM	6929 6930	CD1 ILE C ILE	148	107. 104 80. 131 33. 517 1. 00 34. 57 B	Ċ
ATOM	6931	0 ILE	148 148	103.069 82.641 31.709 1.00 34.54 B 102.492 83.155 32.664 1.00 35.51 B	C
ATOM	6932	N PRO	149	102. 492 83. 155 32. 664 1. 00 35. 51 B 102. 401 82. 199 30. 631 1. 00 32. 42 B	O N
ATOM	6933	CD PRO	149	102. 929 81. 387 29. 525 1. 00 30. 91 B	C
ATOM	6934	CA PRO	149	100. 942 82. 321 30. 526 1. 00 31. 27 B	č
ATOM	6935	CB PRO	149	100. 632 81. 762 29. 134 1. 00 31. 04 B	C C C
ATOM ATOM	6936 6937	CG PRO C PRO	149	101. 963 81. 707 28. 437 1. 00 31. 84 B	C
ATOM	6938	0 PRO	149 149	100. 187 81. 549 31. 592 1. 00 31. 48 B 100. 733 80. 643 32. 221 1. 00 30. 85 B	C
ATOM	6939	N ASN	150	100. 733 80. 643 32. 221 1. 00 30. 85 B 98. 927 81. 919 31. 794 1. 00 31. 40 B	O N
ATOM	6940	CA ASN	150	98. 085 81. 206 32. 744 1. 00 31. 30 B	C
ATOM	6941	CB ASN	150	96. 832 82. 019 33. 108 1. 00 31. 58 B	č
ATOM	6942	CG ASN	150	97. 086 83. 037 34. 211 1. 00 32. 97 B	Č
ATOM ATOM	6943	OD1 ASN	150	97. 676 82. 715 35. 244 1. 00 31. 95 B	0
ATOM	6944 6945	ND2 ASN C ASN	150 150	96. 624 84. 271 34. 004 1. 00 33. 51 B 97. 673 79. 929 32. 013 1. 00 30. 52 B	N
ATOM	6946	0 ASN	150	07 700 70 001	C
ATOM	6947	N ASN	151	97. 722 79. 864 30. 777 1. 00 29. 37 B 97. 269 78. 917 32. 768 1. 00 30. 16 B	O N
ATOM	6948	CA ASN	151	96. 859 77. 657 32. 170 1. 00 29. 53 B	C
ATOM	6949	CB ASN	151	95. 715 77. 881 31. 186 1. 00 33. 04 B	Č
ATOM	6950	CG ASN	151	94. 489 78. 474 31. 850 1. 00 36. 73 B	č
ATOM	6951	OD1 ASN	151	94. 530 79. 586 32. 376 1. 00 38. 47 B	0
ATOM ATOM	6952 6953	ND2 ASN C ASN	151	93. 389 77. 729 31. 831 1. 00 40. 28 B	Ŋ
ATOM	6954	0 ASN	151 151	98. 023 76. 997 31. 452 1. 00 28. 44 B 97. 856 76. 412 30. 382 1. 00 27. 56 B	C
ATOM	6955	N THR	152	97. 856 76. 412 30. 382 1. 00 27. 56 B 99. 212 77. 111 32. 035 1. 00 26. 08 B	0 N
ATOM	6956	CA THR	152	100. 384 76. 489 31. 452 1. 00 24. 37 B	N C
ATOM	6957	CB THR	152	101.682 77.069 32.046 1.00 25.30 B	Č

			FIG. 4-143	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	6959 CG2 THE 6960 C THE 6961 O THE 6962 N GLN 6963 CA GLN 6964 CB GLN 6965 CG GLN 6966 CD GLN 6967 OE1 GLN 6968 NE2 GLN 6969 C GLN	152 152 153 153 153 153 153 153 153	101. 862 78. 407 31. 566 1. 00 25. 07 102. 882 76. 231 31. 643 1. 00 24. 98 100. 257 75. 012 31. 791 1. 00 22. 65 99. 908 74. 652 32. 912 1. 00 21. 72 100. 531 74. 160 30. 815 1. 00 21. 08 100. 407 72. 730 31. 010 1. 00 20. 22 98. 688 72. 573 29. 166 1. 00 20. 23 98. 577 72. 461 27. 669 1. 00 21. 29 99. 365 73. 054 26. 939 1. 00 24. 47 97. 600 71. 703 27. 200 1. 00 20. 51 101. 650 72. 076 31. 578 1. 00 20. 86 101. 574 70. 996 32. 154 1. 00 22. 44 102. 794 72. 729 31. 422 1. 00 20. 43 104. 043 72. 189 31. 934 1. 00 18. 53 104. 387 70. 868 31. 234 1. 00 18. 53 105. 891 69. 559 32. 955 1. 00 17. 98 105. 891 69. 559 32. 955 1. 00 17. 98 107. 261 69. 232	B C C C C C C C C C C C C C C C C C C C
		_	113.621 75.460 37.492 1.00 22.04	B C

				FIC	3. 4 -	144			(Contin	ued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7008 C 7009 C 7010 C 7011 C 7011 C 7012 C 7013 N 7014 C 7015 C 7016 C 7017 7018 C 7020 C 7021 C 7022 C 7023 C 7024 C 7025 C 7026 N 7027 C 7028 C 7029 C 7030 C 7031 C 7032 C 7033 N 7034 C 7035 C 7036 C 7037 N 7038 C 7039 C 7031 C 7032 C 7033 C 7033 C 7034 C 7035 C 7036 C 7037 N 7038 C 7037 N 7038 C 7039 C 7041 C 7041 C 7042 N 7043 C 7044	TREATER TO TREATE TO TREAT	P 157 P 157 P 157 P 157 P 157 R 158 R 158 R 158 R 158 R 159 159 160 160 160 160 160 160 161 161 162 162 162 162 163 163 163 163 163 163 163	113. 193 113. 317 114. 445 113. 779 115. 096 114. 789 116. 198 117. 154 118. 104 118. 550 117. 898 117. 800 118. 641 118. 927 119. 362 120. 041 119. 230 120. 384 120. 598 121. 014 122. 031 123. 383 124. 421 123. 844 121. 606 120. 889 122. 043 121. 706 120. 289 119. 839 119. 584 118. 222 118. 214 119. 019 118. 664 120. 289 119. 839 119. 584 118. 222 118. 214 119. 019 118. 664 120. 824 119. 804 117. 384 116. 730 117. 406 116. 575 117. 113 118. 367 118. 797 120. 103 120. 616 115. 215	76. 650 79. 051 79. 299 79. 846 73. 640 72. 483 74. 211 73. 441 74. 377 75. 444 72. 667 71. 096 70. 860 69. 744 69. 660 71. 738 71. 619 73. 517 73. 272 74. 249 71. 840 74. 952 75. 224 75. 866 77. 266 77. 645 77. 359 78. 296 78. 721 79. 959 81. 094 82. 148 81. 208 82. 283 82. 871 79. 959 81. 094 82. 148 81. 208 82. 283 82. 871 79. 959 81. 094 82. 148 87. 407 77. 578 77. 104	38. 030 37. 286 35. 156 36. 270 33. 153 32. 882 32. 697 31. 996 33. 017 34. 198 32. 650 31. 307 32. 886 31. 599 34. 391 35. 589 34. 391 35. 589 34. 340 35. 589 34. 745 34. 944 36. 053 34. 944 36. 053 34. 944 36. 053 37. 38. 627 38. 629 38. 629 38	1. 00 22. 01 1. 00 22. 77 1. 00 22. 58 1. 00 21. 74 1. 00 22. 79 1. 00 23. 16 1. 00 21. 93 1. 00 22. 68 1. 00 23. 20 1. 00 23. 12 1. 00 23. 12 1. 00 23. 12 1. 00 23. 69 1. 00 24. 45 1. 00 25. 41 1. 00 26. 39 1. 00 27. 71 1. 00 29. 28 1. 00 30. 65 1. 00 33. 70 1. 00 29. 28 1. 00 30. 65 1. 00 33. 70 1. 00 29. 28 1. 00 29. 28 1. 00 30. 65 1. 00 30. 93 1. 00 29. 74 1. 00 30. 93 1. 00 29. 74 1. 00 30. 93 1. 00 29. 32 1. 00 28. 43 1. 00 29. 32 1. 00 28. 43 1. 00 29. 32 1. 00 29. 32 1. 00 20. 70 1. 00 29. 95 1. 00 30. 77 1. 00 29. 24 1. 00 30. 20 1. 00 29. 24 1. 00 30. 20 1. 00 29. 24 1. 00 30. 20 1. 00 29. 25 1. 00 26. 53 1. 00 27 1. 00 29. 26 1. 00 29. 26 1. 00 29. 26 1. 00 29. 26 1. 00 29. 26 1. 00 29. 26 1. 00 21. 68 1. 00 24. 68 1. 00 24. 17 1. 00 22. 69 1. 00 23. 67 1. 00 24. 69	B B B B B B B B B B B B B B B B B B B	NCCCCONCCOCONCCCCCONCCCCONCCCONCOCCCNCNCONCCCCCNCO	

	•							*		
	•				T2 T (٦ ،	1 1 6			(Continued)
					rı	G. 4-	140			
ATOM	7105	CG	ASN	169	96. 682	77. 615	24.662	1.00 27.95	В	С
ATOM	7106		ASN	169	96. 240	78. 640	24. 150	1.00 27.95	В	Ö
ATOM	7107		ASN	169	96. 570	77. 361	25. 961	1.00 32.00	В	N
ATOM	7108	C	ASN	169	98. 463	74. 345	23. 655	1.00 30.33	В	C
ATOM	7109	0	ASN	169	98. 455	74. 541	22.441	1.00 21.23	В	Ö
ATOM	7110	N	ASN	170	99. 031	73. 283	24. 221	1.00 22.01	В	N
ATOM	7111	CA	ASN	170	99. 661	72. 208	23. 459	1.00 20.00	В	
ATOM	7112	CB	ASN	170	98. 615	71.515	22. 592	1.00 18.68	В	C C
ATOM	7113	CG	ASN	170	97. 629	70. 741	23. 412	1.00 18.15	В	Č
ATOM	7114		ASN	170	97. 158	71. 224	24. 440	1.00 16.13	В	Ö
ATOM	7115		ASN	170	97. 300	69. 529	22. 966	1.00 18.92	В	N
ATOM	7116	C	ASN	170	100. 859	72. 581	22. 598	1.00 10.32	В	Č
ATOM	7117	Õ	ASN	170	101. 194	71.861	21.659	1.00 20.36	В	Ö
ATOM	7118	N	ASP	171	101. 504	73. 697	22. 916	1.00 20.30	В	N
ATOM	7119	CA	ASP	171	102. 671	74. 122	22. 160	1.00 23.35	В	C
ATOM	7120	CB	ASP	171	102. 354	75. 364	21.334	1.00 23.05	В	č
ATOM	7121	CG	ASP	171	101.794	75.017	19.978	1.00 23.72	В	Č
ATOM	7122	0D1		171	102. 505	74. 338	19. 210	1.00 23.33	В	Ő
ATOM	7123		ASP	171	100.650	75.415	19.679	1.00 26.97	В	0
ATOM	7124	C	ASP	171	103.850	74. 380	23.073	1.00 23.59	В	Č
ATOM	7125	ŏ	ASP	171	103.672	74.647	24. 264	1.00 24.18	В	ŏ
ATOM	7126	Ň	ILE	172	105.051	74. 301	22. 508	1.00 23.60	. B	Ň ·
ATOM	7127	CA	ILE	172	106. 273	74. 497	23. 281	1.00 25.23	B	Č
ATOM	7128	CB	ILE	172	107. 353	73. 456	22. 885	1.00 23.64	B	č
ATOM	7129		ILE	172	108. 480	73.466	23. 896	1.00 23.11	В	č
ATOM	7130		ILE	172	106. 743	72.056	22.846	1.00 23.95	В	č
ATOM	7131	CD1		172	107.707	70.986	22. 374	1.00 23.66	В	č
ATOM	7132	C	ILE	172	106.878	75.892	23. 129	1.00 25.59	B	Č
ATOM	7133	0	ILE	172	106.881	76.474	22.048	1.00 25.83	B	Ŏ
ATOM	7134	N	TYR	173	107.389	76.414	24. 236	1.00 26.85	B	N .
ATOM	7135	CA	TYR	173	108.025	77.720	24. 272	1.00 27.95	$\tilde{\mathtt{B}}$	Ĉ
ATOM	7136	CB	TYR	173	107.111	78.760	24.933	1.00 27.81	B	Č
ATOM	7137	CG	TYR	173	105.822	79.002	24.190	1.00 29.53	B	Č
ATOM	7138	CD1	TYR	173	104.788	78.063	24.226	1.00 29.72	В	Č
ATOM	7139	CE1	TYR	173	103.599	78. 271	23.535	1.00 29.08	. B	Č
ATOM	7140	CD2		173	105.634	80.162	23.439	1.00 28.71	В	Č
ATOM	7141		TYR	173	104. 444	80.381	22.740	1.00 30.14	В	C
ATOM	7142	CZ	TYR	173	103.432	79.429	22.794	1.00 30.82	В	C
ATOM	7143	OH	TYR	173	102. 258	79.625	22.103	1.00 31.14	В	0
ATOM	7144	C	TYR	173	109.308	77.592	25.080	1.00 28.66	В	C
ATOM	7145	0	TYR	173	109.412	76.735	25.960	1.00 28.10	В	0
ATOM	7146	N	VAL	174	110.276	78.451	24.782	1.00 29.35	В	N
ATOM	7147	CA	VAL	174	111.551	78.443	25.480	1.00 29.22	В	Ċ
ATOM	7148	CB	VÁL	174	112.669	77.855	24.587	1.00 29.66	В	Č ·
ATOM	7149	CG1		174	114.006	77.936	25.303	1.00 30.07	В	Ċ
ATOM	7150		VAL	174	112.351	76.403	24.231	1.00 30.25	В	C .
ATOM	7151	C	VAL	174	111.953	79.857	25.887	1.00 30.16	В	C
ATOM	7152	0	VAL	174	111.787	80.804	25.125	1.00 31.81	В	0
ATOM	7153	N	LYS	175	112.474	79.990	27. 099	1.00 29.78	В	N

					FΙ	G. 4	-147	7		(Con	tinued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7154 7155 7156 7157 7158 7159 7160 7161 7162 7163 7164 7165 7166 7167 7167 7172 7173 7174 7175 7176 7177 7178 7178 7180 7181 7182 7183 7184 7185 7186 7187 7187 7188 7189 7191 7192 7193 7194 7195 7197 7197 7198 7199 7200 7201 7202	CBCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	LYS	175 175 175 175 175 175 175	F I 112. 940 112. 090 110. 809 109. 876 110. 664 114. 382 114. 662 115. 294 116. 710 117. 572 118. 942 117. 697 116. 985 116. 182 117. 910 116. 085 116. 182 119. 324 120. 511 118. 914 114. 762 113. 905 114. 495 115. 451 113. 160 113. 383 114. 862 112. 451 113. 198 112. 560 113. 383 114. 454 115. 419 114. 457 115. 419 114. 457 115. 621 116. 828 115. 522 111. 898 111. 406 111. 462	81. 269 81. 725 82. 428 82. 551 83. 384 84. 791 81. 107 80. 355 81. 813 81. 764 82. 363 82. 730 81. 354 80. 941 82. 528 82. 251 83. 489 84. 296 85. 611 85. 440	27. 608 28. 794 28. 413 29. 611 30. 725 30. 307 28. 064 28. 999 27. 401 27. 749 26. 624 27. 146 25. 483 24. 861 29. 044 29. 768 29. 330 30. 543 30. 241 29. 770 30. 916	1.00 28.47 1.00 28.38 1.00 29.46 1.00 32.27 1.00 31.57 1.00 28.80 1.00 28.36 1.00 28.58 1.00 28.19 1.00 27.21 1.00 25.54 1.00 28.29 1.00 27.38 1.00 29.36	B B B B B B B B B B B B B B B B B B B	CCCCCNCONCCCCCONCCCCONCCCCCONCCCONCONCCCON	
				•	LIDATITLITE	- 011555					

			FIC 4 140	(Continued)
			FIG. 4-148	
ATOM ATOM	7203 CD PR 7204 CA PR		111. 853 88. 984 25. 784 1. 00 33. 21 B	
ATOM	7205 CB PR		110 007 07 000	C
ATOM	7206 CG PR		110 001 00 000	C
ATOM	7207 C PR		110. 681 85. 608 25. 397 1. 00 33. 21 B	C C
ATOM	7208 O PR		111. 829 85. 180 25. 497 1. 00 33. 18 B	0
ATOM	7209 N SE		109. 654 84. 838 25. 070 1. 00 33. 87 B	N N
ATOM	7210 CA SE		109. 835 83. 415 24. 829 1. 00 32. 06 B	Č
ATOM	7211 CB SE		108. 752 82. 622 25. 547 1. 00 31. 33 B	č
ATOM	7212 OG SE	_	107. 505 82. 817 24. 909 1. 00 30. 50 B	0
ATOM ATOM	7213 C SE 7214 O SE		109. 759 83. 117 23. 350 1. 00 31. 89 B	C
ATOM	7214 O SE 7215 N TY		109.077 83.812 22.606 1.00 33.14 B	0
ATOM	7216 CA TY		110. 463 82. 077 22. 927 1. 00 31. 53 B 110. 453 81. 677 21. 532 1. 00 30. 47 B	N
ATOM	7217 CB TY		111 000 01 150	C
ATOM	7218 CG TY		110 000 00 115	C
ATOM	7219 CD1 TY		112. 962 82. 117 21. 408 1. 00 32. 75 B 113. 490 82. 235 22. 696 1. 00 32. 39 B	C
ATOM	7220 CE1 TY		114. 517 83. 134 22. 977 1. 00 33. 30 B	Č
ATOM	7221 CD2 TYI		113. 492 82. 926 20. 398 1. 00 33. 06 B	č
ATOM	7222 CE2 TYI		114. 520 83. 832 20. 667 1. 00 34. 20 B	č
ATOM ATOM	7223 CZ TYF 7224 OH TYF		115. 028 83. 932 21. 959 1. 00 34. 92 B	Ċ
ATOM	7224 OH TYF 7225 C TYF		116. 036 84. 832 22. 233 1. 00 34. 60 B	0
ATOM	7226 0 TYF		109. 423 80. 568 21. 384 1. 00 29. 28 B 109. 387 79. 645 22. 196 1. 00 29. 66 B	C
ATOM	7227 N ARG		100 570 00 050 00 000	0
ATOM	7228 CA ARG		107 570 70 001	N
ATOM	7229 CB ARG		107. 573 79. 531 20. 148 1. 00 26. 57 B	C
ATOM	7230 CG ARG		105. 215 79. 191 19. 285 1. 00 28. 64 B	C C
ATOM	7231 CD ARG		103.860 79.825 19.004 1.00 30.29 B	Č
ATOM	7232 NE ARG		102. 827 78. 805 18. 831 1. 00 31. 47 B	Ň
ATOM ATOM	7233 CZ ARG 7234 NH1 ARG		101. 526 79. 052 18. 706 1. 00 29. 99 B	Č
ATOM	7235 NH2 ARG		100. 678 78. 048 18. 552 1. 00 30. 76 B	N
ATOM	7236 C ARG		101.068 80.294 18.740 1.00 30.05 B 108.185 78.553 19.272 1.00 26.51 B	N
ATOM	7237 0 ARG		100 275 70 754 10 255	C
ATOM	7238 N ILE	185	100 400 77 411 10 070	0
ATOM	7239 CA ILE	185	108. 493 77. 411 19. 876 1. 00 24. 50 B 109. 112 76. 303 19. 165 1. 00 22. 88 B	N
MOTA	7240 CB ILE	185	109. 773 75. 319 20. 159 1. 00 23. 12 B	C C
ATOM	7241 CG2 ILE	185	110. 492 74. 216 19. 405 1. 00 22. 56 B	Č
ATOM	7242 CG1 ILE	185	110. 753 76. 067 21. 064 1. 00 22. 32 B	č
ATOM ATOM	7243 CD1 ILE 7244 C ILE	185	111. 869 76. 770 20. 324 1. 00 21. 93 B	Č
ATOM	7244 C ILE 7245 O ILE	185 185	108. 148 75. 516 18. 275 1. 00 24. 00 B	. C
ATOM	7246 N THR	186	108. 569 74. 930 17. 275 1. 00 25. 07 B 106. 866 75. 489 18. 632 1. 00 22. 70 B	0
ATOM	7247 CA THR	186	105 000 54 550	N
ATOM	7248 CB THR	186	100 400 70 440	C
ATOM	7249 OG1 THR	186	105. 490 73. 440 18. 541 1. 00 22. 83 B 105. 058 73. 727 19. 877 1. 00 27. 42 B	C
ATOM	7250 CG2 THR	186	106. 665 72. 491 18. 595 1. 00 19. 86 B	0 C
ATOM	7251 C THR	186	104. 620 75. 548 17. 537 1. 00 23. 45 B	Č

				FIG. 4-150	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7302 C 7303 C 7304 N 7305 C 7306 C 7307 C 7308 C 7310 C 7311 C 7312 N 7313 C 7314 C 7315 C 7316 C 7317 C 7318 C 7317 C 7318 C 7317 C 7318 C 7319 C 7320 N 7321 C 7322 C 7323 C 7324 C 7325 C 7326 C 7327 C 7328 C 7329 C 7329 C 7330 C 7331 C	O ILE N ILE CA ILE CB ILE CG2 ILE CG1 ILE CD1 ILE TYR CA TYR CB T	192 192 193 193 193 193 193 193 193 194 194 194 194 195 195 195 195 195 195 195 195	89. 671 67. 908 23. 548 1. 00 32. 44 B 93. 072 66. 329 20. 602 1. 00 25. 95 B 92. 431 65. 426 20. 065 1. 00 27. 81 B 94. 091 66. 926 20. 000 1. 00 25. 46 B 94. 485 66. 512 18. 665 1. 00 25. 50 B 93. 970 67. 502 17. 595 1. 00 26. 97 B 94. 426 67. 057 16. 212 1. 00 26. 11 B 92. 441 67. 552 17. 621 1. 00 27. 90 B 91. 784 66. 246 17. 210 1. 00 29. 23 B 95. 994 66. 390 18. 546 1. 00 25. 04 B 96. 519 65. 297 18. 334 1. 00 26. 34 B 96. 691 67. 510 18. 682 1. 00 21. 47 B 98. 139 67. 505 18. 589 1. 00 21. 47 B 98. 618 68. 429 17. 456 1. 00 2	(Continued) 0 C 0 N C C C C C C C C C C C C C C C
ATOM ATOM	7333 CA 7334 CE	B ASN	196 196	103. 762 68. 941 21. 725 1. 00 17. 79 B 104. 011 70. 187 20. 867 1. 00 17. 21 B	C C
ATOM ATOM ATOM	7337 NI	G ASN D1 ASN D2 ASN	196 196 196	103. 366 70. 106 19. 489 1. 00 17. 04 B 103. 769 69. 311 18. 632 1. 00 16. 41 B 102. 362 70. 943 19. 267 1. 00 17. 01 B	C O N
ATOM ATOM ATOM	7338 C 7339 O 7340 N	ASN ASN GLY	196 196 197	104. 380 69. 160 23. 104 1. 00 18. 89 B 103. 976 70. 066 23. 828 1. 00 21. 80 B	O
ATOM ATOM	7341 CA 7342 C	A GLY GLY	197 197	105. 355 68. 344 23. 479 1. 00 18. 21 B 105. 976 68. 533 24. 778 1. 00 18. 42 B 105. 185 67. 948 25. 941 1. 00 18. 43 B	N C C
ATOM ATOM ATOM	7343 O 7344 N 7345 CA		197 198 198	105. 660 67. 954 27. 088 1. 00 17. 86 B 103. 976 67. 469 25. 654 1. 00 15. 16 B 103. 129 66. 842 26. 667 1. 00 14. 58 B	O N C
ATOM ATOM ATOM ATOM	7348 CG	2 ILE 1 ILE	198 198 198	101. 956 67. 740 27. 160 1. 00 12. 66 B 102. 477 68. 784 28. 109 1. 00 10. 73 B 101. 189 68. 334 25. 970 1. 00 14. 13 B	C C C
UTON	1040 UD	1 ILE	198	99. 936 69. 129 26. 368 1. 00 13. 46 B	С

					TOTO A 1E1	(Continued)				
					FIG. 4-151	•				
ATOM ATOM	7350 7351		ILE		102. 523 65. 585 26. 101 1. 00 14. 46 B	С				
ATOM	7352		ILE THR		102. 354 65. 447 24. 895 1. 00 16. 78 B 102. 182 64. 671 26. 990 1. 00 15. 77 B	0				
ATOM	7353				10: 000 00 000	N				
ATOM	7354				101 000 10 100	C				
ATOM	7355				104 000 00 00	C				
ATOM	7356			199	101. 683 62. 861 28. 937 1. 00 12. 99 B 103. 473 62. 043 27. 534 1. 00 15. 54 B	0 C				
ATOM	7357	C	THR	199	100.085 63.448 26.522 1.00 15.87 B	C				
ATOM	7358		THR	199	99. 452 64. 311 27. 133 1. 00 16. 77 B	Ö				
ATOM	7359		ASP	200	99. 510 62. 534 25. 745 1. 00 16. 29 B	Ň				
ATOM	7360			200	98. 058 62. 450 25. 619 1. 00 16. 42 B	Ĉ				
ATOM	7361	CB		200	97. 654 61. 812 24. 279 1. 00 17. 56 B	C				
ATOM ATOM	7362			200	97. 960 60. 321 24. 207 1. 00 19. 40 B	C				
ATOM	7363 7364		1 ASP 2 ASP	200 200	98. 894 59. 847 24. 892 1. 00 20. 07 B	0				
ATOM	7365	C	ASP	200	97. 267 59. 624 23. 438 1. 00 19. 79 B 97. 657 61. 578 26. 806 1. 00 15. 56 B	0				
ATOM	7366	ő	ASP	200	00 500 44 050	C				
ATOM	7367	N	TRP	201	00 101 01 151	0				
ATOM	7368	CA	TRP	201	96. 404 61. 151 26. 889 1. 00 14. 09 B 96. 003 60. 368 28. 049 1. 00 13. 08 B	N C				
ATOM	7369	CB	TRP	201	94. 503 60. 106 28. 037 1. 00 13. 25 B	Č				
ATOM	7370	· CG	TRP	201	94. 023 59. 554 29. 348 1. 00 12. 63 B	Č				
ATOM	7371		TRP	201	94. 135 58. 198 29. 801 1. 00 10. 35 B	č				
ATOM	7372		TRP	201	93. 610 58. 150 31. 110 1. 00 11. 08 B	Č				
ATOM ATOM	7373 7374		TRP	201	94. 634 57. 020 29. 228 1. 00 8. 52 B	C				
ATOM	7375		TRP TRP	201	93. 449 60. 253 30. 370 1. 00 12. 43 B	C				
ATOM	7376		TRP	201 201	93. 198 59. 416 31. 434 1. 00 12. 21 B	N				
ATOM	7377		TRP	201	93. 567 56. 967 31. 858 1. 00 11. 85 B 94. 596 55. 847 29. 968 1. 00 8 91 B	C				
ATOM	7378		TRP	201	D4 005 55 000	C				
ATOM	7379	C	TRP	201	00 710 70 040 00	C				
ATOM	7380	0	TRP	201	96. 719 59. 040 28. 264 1. 00 14. 63 B 97. 197 58. 766 29. 366 1. 00 14. 84 B	C 0				
ATOM	7381	N	ŸAL	202	96. 795 58. 213 27. 224 1. 00 14. 84 B	N N				
ATOM	7382	CA	VAL	202	97. 413 56. 902 27. 369 1. 00 13. 74 B	Č				
ATOM	7383	CB	VAL	202	97. 028 55. 966 26. 190 1. 00 11. 30 B	č ·				
ATOM	7384	CGI	VAL	202	97. 960 56. 155 25. 010 1. 00 8. 57 B	Č				
ATOM ATOM	7385 7386	C	VAL	202	97. 028 54. 541 26. 667 1. 00 8. 82 B	C				
ATOM	7387	Ö	VAL VAL	202 202	98. 929 56. 920 27. 556 1. 00 15. 45 B	C				
ATOM	7388	N	TYR	202	99. 471 56. 095 28. 292 1. 00 16. 05 B 99. 616 57. 857 26. 906 1. 00 15 45 B	0				
ATOM	7389	CA	TYR	203	- 2, 000 10, 10 D	N				
ATOM	7390	CB	TYR	203	101 000 000	C				
ATOM	7391	CG	TYR	203	101. 656 58. 918 26. 035 1. 00 12. 37 B 102. 248 58. 238 24. 823 1. 00 8. 90 B	C				
ATOM	7392	CD1		203	101. 461 57. 938 23. 709 1. 00 8. 82 B	C C				
ATOM	7393	CE1		203	101. 989 57. 260 22. 619 1. 00 7. 48 B	C				
ATOM	7394	CD2		203	103. 587 57. 844 24. 812 1. 00 5. 53 B	č				
ATOM	7395	CE2		203	104. 128 57. 167 23. 727 1. 00 6. 51 B	C C C				
ATOM ATOM	7396 7397	CZ OH	TYR	203	103. 325 56. 874 22. 634 1. 00 8. 49 B					
ATOM	7398	C	TYR TYR	203 203	103.849 56.175 21.572 1.00 8.01 B	0				
a a.a. Wald	1000	U	111/		101. 438 58. 371 28. 471 1. 00 13. 68 B	С				
	SUBSTITUTE SHEET (RULE 26)									

										(Con	tinued)
		4			FIC	3.4-	1 5 2	•			
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7399 7400 7401 7402 7403 7404 7405 7406	O N CA CB CG CD OE1	TYR GLU GLU GLU GLU GLU GLU	203 204 204 204 204 204 204 204	102. 369 100. 706 100. 963 99. 975 100. 174 98. 950 98. 197 98. 753	57. 832 59. 335 59. 827 60. 936 61. 457 62. 154 62. 785 62. 085	29. 056 29. 020 30. 376 30. 743 32. 161 32. 731 31. 964 33. 962	1. 00 12. 65 1. 00 15. 26 1. 00 16. 69 1. 00 16. 67 1. 00 17. 47 1. 00 17. 71 1. 00 19. 00 1. 00 18. 59	B B B B B B	0 N C C C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7407 7408 7409 7410 7411 7412 7413 7414	C O N CA CB CG CD OE1		204 204 205 205 205 205 205 205 205	100. 831 101. 681 99. 745 99. 442 97. 925 97. 453 97. 414 97. 038	58. 740 58. 597 57. 980 56. 932 56. 727 55. 436 55. 494 54. 466	31. 437 32. 305 31. 353 32. 315 32. 344 32. 995 34. 515 35. 118	1. 00 17. 37 1. 00 18. 22 1. 00 18. 89 1. 00 19. 55 1. 00 20. 80 1. 00 23. 74 1. 00 26. 68 1. 00 28. 71	B B B B B B	C O N C C C C C O	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7415 7416 7417 7418 7419 7420 7421	C O N CA CB CG	GLU GLU GLU GLU GLU GLU	205 205 205 206 206 206 206	97. 744 100. 132 100. 525 100. 291 100. 876 99. 989 98. 535	56. 547 55. 578 54. 957 55. 124 53. 808 53. 016 52. 921	35. 106 32. 131 33. 107 30. 893 30. 660 29. 705 30. 139	1.00 26.12 1.00 19.27 1.00 19.31 1.00 18.93 1.00 18.63 1.00 18.05 1.00 20.39	B B B B B	0 C O N C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7422 7423 7424 7425 7426 7427 7428 7429	CD OE1 OE2 C O N CA CB	GLU GLU GLU GLU VAL VAL VAL	206 206 206 206 206 207 207	98. 359 97. 205 99. 375 102. 293 102. 976 102. 744 104. 092	52. 143 51. 905 51. 768 53. 766 52. 761 54. 844 54. 855	31. 422 31. 821 32. 037 30. 136 30. 292 29. 509 28. 968	1.00 20.74 1.00 21.45 1.00 22.90 1.00 19.32 1.00 20.01 1.00 20.90 1.00 20.95	B B B B	C 0 0 C 0 N C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7430 7431 7432 7433 7434 7435 7436	CG1	VAL VAL VAL VAL PHE PHE PHE	207 207 207 207 207 208 208 208	104. 101 105. 486 103. 048 105. 080 106. 052 104. 833 105. 743 105. 877	55. 347 55. 151 54. 592 55. 691 55. 160 56. 989 57. 870 59. 201	27. 509 26. 918 26. 684 29. 775 30. 301 29. 888 30. 611 29. 863	1.00 21.52 1.00 22.17 1.00 19.10 1.00 21.67 1.00 25.32 1.00 21.55 1.00 21.33 1.00 21.28	B B B B B B	C C C C O N C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7437 7438 7439 7440 7441 7442 7443	CG CD1 CD2 CE1	PHE PHE PHE PHE PHE PHE PHE	208 208 208 208 208 208 208 208	105. 877 106. 571 107. 890 105. 893 108. 525 106. 521 107. 837 105. 444		29. 803 28. 536 28. 464 27. 353 27. 230 26. 109 26. 048 32. 082	1.00 21.28 1.00 21.92 1.00 20.63 1.00 22.58 1.00 22.52 1.00 22.24 1.00 22.76 1.00 21.89	B B B B B B	0000000	
ATOM ATOM ATOM ATOM	7444 7445 7446 7447	O N CA CB	PHE SER SER SER	208 209 209 209	106. 298 104. 261 103. 922 104. 689	58. 727 57. 811 58. 094 57. 165	32. 768 32. 577 33. 976 34. 905	1.00 21.89 1.00 23.07 1.00 20.48 1.00 19.86 1.00 18.09	В В В В	C O N C C	

					(Continued)
				FIG. 4-153	
ATOM ATOM	7443			104.383 55.820 34.601 1.00 21.42 B	0
ATOM	7449 7450			104. 285 59. 543 34. 286 1. 00 20. 55 B	C
ATOM	745			104. 780 59. 877 35. 367 1. 00 19. 53 B	0
ATOM	7452			104.031 60.394 33.302 1.00 20.69 B	N
ATOM	7453			104. 319 61. 809 33. 393 1. 00 20. 47 B	C
ATOM	7454			105. 809 62. 044 33. 228 1. 00 20. 63 B 103. 545 62. 492 32. 275 1. 00 20. 53 B	C
ATOM	7455			10 200 201 00 B	C
ATOM	7456				0
ATOM	7457			100 800	N
ATOM	7458			102. 733 64. 634 31. 390 1. 00 20. 95 B 101. 944 65. 681 32. 175 1. 00 18. 35 B	C C
ATOM	7459			100.984 66.566 31.411 1.00 15.38 B	C
ATOM	7460			100. 257 66. 086 30. 324 1. 00 14. 13 B	C
ATOM	7461			99. 310 66. 879 29. 694 1. 00 12. 47 B	C
ATOM	7462		211	100. 738 67. 863 31. 846 1. 00 11. 95 B	Č
ATOM	7463			99. 799 68. 657 31. 231 1. 00 12. 21 B	č
ATOM	7464			99. 087 68. 165 30. 156 1. 00 13. 68 B	č
ATOM	7465			98. 158 68. 977 29. 550 1. 00 12. 73 B	<u>0</u> ·
ATOM	7466			103. 781 65. 283 30. 508 1. 00 22. 11 B	Č
ATOM	7467	0 TYR		103. 512 65. 742 29. 406 1. 00 23. 55 B	0
ATOM ATOM	7468	N SER	212	105. 000 65. 294 31. 017 1. 00 23. 17 B	N
ATOM	7469 7470	CA SER CB SER	212	106. 112 65. 877 30. 310 1. 00 22. 03 B	C
ATOM	7471	CB SER OG SER	212	107. 286 66. 055 31. 265 1. 00 22. 38 B	C
ATOM	7472	C SER	212 212	108.441 66.477 30.567 1.00 24.83 B	0
ATOM	7473	0 SER	212	106. 547 65. 017 29. 141 1. 00 22. 20 B 106. 651 63. 802 29. 256 1. 00 22. 93 B	C
ATOM	7474	N ALA	213	100 701 07 000	0
ATOM	7475	CA ALA	213	107 007 05 011	N
ATOM	7476	CB ALA	213	100 155 01 000	C
ATOM	7477	C ALA	213	106. 157 64. 882 25. 803 1. 00 19. 85 B 108. 360 65. 942 26. 301 1. 00 21. 17 B	C C
ATOM	7478	0 ALA	213	108. 443 66. 254 25. 109 1. 00 20. 14 B	0
ATOM	7479	N LEU	214	109. 175 66. 409 27. 243 1. 00 21. 21 B	N N
ATOM	7480	CA LEU	214	110. 298 67. 295 26. 961 1. 00 22. 06 B	Č
ATOM	7481	CB LEU	214	110.049 68.697 27.534 1.00 21.02 B	č
ATOM	7482	CG LEU	214	108.958 69.546 26.878 1.00 20.19 B	č
ATOM	7483	CD1 LEU	214	108.840 70.872 27.603 1.00 21.72 B	Č
ATOM ATOM	7484 7485	CD2 LEU	214	109. 292 69. 779 25. 426 1. 00 22. 01 B	C
ATOM	7486	C LEU O LEU	214	111. 528 66. 688 27. 615 1. 00 22. 30 B	C
ATOM	7487	O LEU N TRP	214	111. 442 66. 131 28. 703 1. 00 25. 61 B	0 ·
ATOM	7488	CA TRP	215 215	112. 674 66. 795 26. 957 1. 00 21. 71 B	· N
ATOM	7489	CB TRP	215	113. 904 66. 237 27. 497 1. 00 19. 34 B	C
ATOM	7490	CG TRP	215	114. 112 64. 833 26. 942 1. 00 18. 71 B 113. 018 63. 863 27. 294 1. 00 18. 43 B	C
ATOM	7491	CD2 TRP	215	111 010 00 101 00 100	C
ATOM	7492	CE2 TRP	215	111 150 00 000	C
ATOM	7493	CE3 TRP	215	111 100 00 015 05 100	C
ATOM	7494	CD1 TRP	215	111. 482 63. 845 25. 186 1. 00 17. 01 B 112. 890 63. 155 28. 456 1. 00 15. 04 B	C
ATOM	7495	NE1 TRP	215	111. 781 62. 356 28. 400. 1. 00 13. 49 B	N C·
ATOM	7496	CZ2 TRP	215	109. 996 61. 949 26. 682 1. 00 14. 75 B	N C
				OUDOTHUR OF THE STATE OF THE ST	U

					DIC		4			(Continue	ed)
					FIG.	4 -	155				
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7546 7547 7548 7549 7550 7551 7552 7553 7555 7556 7557 7558 7559 7560 7561	CD: CE: CZ CZ C O N CA CB CG	THE PHE PHE PHE LEU LEU LEU LEU LEU LEU LEU LEU LEU LE	222 222 222 222 222 222 222 222 222 22	119. 862 7 120. 305 7 119. 158 7 119. 480 6 120. 722 6 121. 955 6 120. 661 7 123. 115 6 121. 815 7 123. 046 6 117. 949 7 118. 066 7 116. 780 7 115. 540 7 114. 618 7 113. 248 7	3. 551 1. 386 0. 921 9. 645 9. 723 9. 384 0. 111 9. 425 0. 158 9. 814 0. 282 0. 746 0. 442 1. 667 1. 340	23. 201 23. 619 22. 850 22. 069 21. 246 21. 797 19. 912 21. 031 19. 693 23. 723 24. 901 23. 119 23. 789 23. 878 24. 503	1. 00 28. 29 1. 00 25. 34 1. 00 25. 13 1. 00 25. 65 1. 00 26. 36 1. 00 26. 35 1. 00 25. 81 1. 00 28. 19 1. 00 24. 55 1. 00 24. 38 1. 00 24. 38 1. 00 22. 85 1. 00 21. 81 1. 00 20. 49	B B B B B B B B B B B B B B B B B B B	0 N C C C C C C C C C C C C C C C C C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7562 7563 7564 7565 7566 7567 7568 7569 7570 7571 7572 7573 7574 7575 7576	CD2 C O N CA CB C O N CA CB CCA CB CCA CB	LEU LEU ALA ALA ALA ALA ALA TYR TYR TYR TYR	223 223 223 224 224 224 224 225 225 225 225 225	113. 469 70 112. 389 73 114. 885 69 114. 462 69 114. 834 68 114. 201 67 114. 935 69 112. 761 66 112. 498 67 111. 825 66 110. 423 66 109. 733 67 109. 648 68 110. 680 69	0. 684 2. 587 9. 380 9. 650 3. 162 7. 062 5. 776 5. 968 7. 111 6. 755 6. 635 7. 997 8. 624	25. 860 24. 644 22. 934 21. 808 23. 459 22. 753 23. 038 23. 248 24. 444 22. 328 22. 703 22. 701 21. 332 20. 849	1. 00 21. 10 1. 00 18. 49 1. 00 23. 23 1. 00 22. 62 1. 00 23. 47 1. 00 23. 08 1. 00 24. 27 1. 00 23. 38 1. 00 23. 37 1. 00 23. 10 1. 00 21. 31 1. 00 18. 23 1. 00 18. 56 1. 00 16. 52	B B B B B B B B B B B B B B B B B B B	C C C O N C C C C C C C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7577 7578 7579 7580 7581 7582 7583 7584 7585 7586 7587 7588 7589 7590 7591 7592 7593	CD2		225 225 225 225 225 225 226 226 226 226	108. 543 68 108. 466 68 109. 502 69 109. 431 70 109. 705 65 110. 143 65 108. 596 65 107. 811 64 107. 485 62 106. 528 64 106. 107 65 105. 912 64 104. 659 64 104. 823 65 103. 512 66 103. 554 66 103. 724 66	. 410 . 968 . 709 . 300	19. 909 19. 457 18. 139 17. 670 16. 249	1. 00 13. 07 1. 00 16. 18 1. 00 14. 89 1. 00 12. 68 1. 00 14. 06 1. 00 21. 55 1. 00 22. 86 1. 00 20. 96 1. 00 19. 66 1. 00 19. 73 1. 00 21. 22 1. 00 16. 70 1. 00 17. 01 1. 00 17. 47 1. 00 18. 65 1. 00 18. 45 1. 00 18. 91	B B B B B B B B B B B B B B B B B B B	C C C C C O N C C C C C C O N C C C C C	

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					F.1 (G. 4-	156			
ATOM	acne	0	OLM	007	100 651	00 041	10.074	1 00 17 01	n	0
ATOM	7595	C	GLN	227	103.651	63. 841	19. 274	1.00 17.21	В	C
ATOM	7596	0	GLN	227	103. 931	62.850	18. 594	1.00 17.76	В	0
ATOM	7597	N	PHE	228	102.483	63. 990	19.888	1.00 16.03	В	N
ATOM	7598	CA	PHE	228	101.447	62. 980	19. 768	1.00 17.64	В	C
ATOM	7599	CB	PHE	228	100.985	62. 524	21.158	1.00 14.78	В	C
ATOM	7600	CG	PHE	228	102.111	62. 105	22.065	1.00 13.03	В	C
ATOM	7601		PHE	228	102.659	63.003	22. 982	1.00 12.33	В	C
ATOM	7602		PHE	228	102.653	60.826	21.978	1.00 12.01	В	C
ATOM	7603		PHE	228	103.732	62.636	23. 796	1.00 9.77	В	C
ATOM	7604		PHE	228	103.725	60.450	22. 786	1.00 11.27	В	C
ATOM ATOM	7605	CZ	PHE	228 228	104. 267	61.360	23.698	1.00 9.50	В	C
ATOM	7606 7607	C	PHE PHE	228	100. 263 99. 894	63. 523 64. 697	18. 955 19. 064	1.00 18.96 1.00 19.98	В	C
ATOM	7608	O N	ASN	228 229	99. 685	62.657	18.133	1.00 19.98	В	0 N
ATOM	7609	CA	ASN	229	98. 548	63. 002	17. 285	1.00 20.11	B B	N C
ATOM	7610	CB	ASN	229	98. 965	62. 867	15.819	1.00 20.74	В	C C
ATOM	7611	CG	ASN	229	97. 980	63. 488	14. 867	1.00 27.56	В	C
ATOM	7612		ASN	229	96. 795	63. 610	15. 174	1.00 21.30	В	0
ATOM	. 7613		ASN	229	98. 467	63. 871	13.692	1.00 30.76	В	N N
ATOM	7614	C	ASN	229	97. 435	61. 995	17.609	1.00 21.10	В	C
ATOM	7615	ŏ	ASN	229	97. 550	60.816	17. 283		В	0
ATOM	7616	N	ASP	230	96. 369	62. 444	18. 260	1.00 22.16	В	N N
ATOM	7617	CA	ASP	230	95. 277	61.534	18.608	1.00 24.31	В	Č
ATOM	7618	CB	ASP	230	94.877	61.683	20.079	1.00 23.86	В	č
ATOM	7619	ĊĠ	ASP	230	95. 999	61.332	21.027	1.00 25.25	В	č
ATOM	7620		ASP	230	95. 701	60.914		1.00 27.89	В	ŏ
ATOM	7621		ASP	230	97. 180	61.485	20.656	1.00 27.78	B	0 .
ATOM	7622	C	ASP	230	94.056	61.776	17.740	1.00 24.83	B	Č
ATOM	7623	0	ASP	230	92.927	61.496	18.148	1.00 24.00	B	Ö
ATOM	7624	N	THR	231	94. 297	62.284	16.536	1.00 25.37	B	Ň
ATOM	7625	CA	THR	231	93. 229	62.582	15.593	1.00 26.24	B	Ċ
ATOM	7626	CB	THR	231	93.802	62.868	14.193	1.00 25.71	B	Č
ATOM	7627	0G1	THR	231	94.439	64.151	14.194	1.00 26.78	В	0
ATOM	7628	CG2	THR	231	92. 702	62.851	13.150	1.00 23.72	В	C
ATOM	7629	C	THR	231	92. 148	61.510	15.467	1.00 27.04	В	C
ATOM	7630	0	THR	231	90.964	61.815	15.604	1.00 29.05	В	0
ATOM	7631	N	GLU	232	92.545	60.265	15. 211	1.00 27.00	В	N
ATOM	7632	CA	GLU	232	91.574	59. 183	15.038	1.00 26.30	В	C
ATOM	7633	CB	GLU	232	92.017	58. 286	13.877	1.00 29.71	В	C
ATOM	7634	CG	GLU	232	92.177	59.036	12.563	1.00 36.71	В	C
ATOM	7635	CD	GLU	232	92. 971	58. 253	11.519	1.00 39.94	В	C
ATOM	7636		GLU	232	92. 434	57. 273	10.943	1.00 41.61	В	0
ATOM	7637		GLU	232	94. 142	58. 623	11. 286	1.00 39.28	В	0
ATOM	7638	C	GLU	232	91.320	58. 328	16. 282	1.00 23.78	В	C
ATOM	7639	0	GLU	232	90. 683	57. 280	16. 208	1.00 23.18	В	0
ATOM	7640	N	VAL	233	91.823	58. 763	17. 427	1.00 21.91	В	N
ATOM	7641	CA	VAL	233	91.608	58.010	18.652	1.00 20.18	В	C
ATOM	7642		VAL	233	92.651	58. 375	19.727	1.00 20.26	В	C
ATOM	7643	UUI	VAL	233	92. 352	57.627	21.016	1.00 18.23	В	С

				(Continued)
			FIG. 4-157	(
ATOM			94. 050 58. 032 19. 223 1. 00 18. 80 B	С
ATOM			90. 218 58. 339 19. 175 1. 00 18. 04 B	Č
ATOM			89. 886 59. 507 19. 378 1. 00 19. 49 B	
ATOM			89. 383 57. 315 19. 394 1. 00 16. 04 B	
ATOM			89. 633 55. 876 19. 231 1. 00 14. 37 B	
ATOM			88. 025 57. 544 19. 896 1. 00 15. 33 B	Č
ATOM			87. 461 56. 133 20. 030 1. 00 13. 91 B	Č ·
ATOM			88. 247 55. 363 19. 013 1. 00 12. 89 B	C
ATOM ATOM			88. 048 58. 275 21. 227 1. 00 14. 45 B	C
ATOM	7653 O . PR		89. 043 58. 242 21. 950 1. 00 13. 13 B	0
ATOM	7654 N LE		86. 941 58. 927 21. 547 1. 00 14. 92 B	N
ATOM	7655 CA LEI 7656 CB LEI		86. 831 59. 676 22. 791 1. 00 13. 91 B	C
ATOM			86. 131 61. 005 22. 536 1. 00 14. 93 B	C
ATOM	7657 CG LEI 7658 CD1 LEI		86. 627 61. 937 21. 434 1. 00 16. 83 B	C
ATOM	7659 CD2 LEU		85. 581 63. 030 21. 198 1. 00 17. 90 B	C
ATOM	7660 C LEU		87. 963 62. 534 21. 833 1. 00 14. 85 B	С
ATOM	7661 O LEI		85. 998 58. 911 23. 803 1. 00 12. 70 B 84. 941 58. 385 23. 456 1. 00 13. 27 B	C
ATOM	7662 N ILE		90 400 50 001	0
ATOM	7663 CA ILE		OF (10 FO 10F CO 10F)	N
ATOM	7664 CB ILE		00 000 50 000	C C C C
ATOM	7665 CG2 ILE		07 916 50 400 07 070	C
ATOM	7666 CG1 ILE		OF 900 FT 040 00 15	C
ATOM	7667 CD1 ILE		04 405 50 100 00 000	C
ATOM	7668 C ILE		94 774 50 000 00 450	C
ATOM	7669 0 ILE	236	0F 977 CO FOO OG 100	C
ATOM	7670 N GLU	237	00 407 50 150 00 500	0 .
ATOM	7671 CA GLU	237	83. 497 59. 156 26. 741 1. 00 13. 69 B 82. 651 60. 267 27. 150 1. 00 14. 30 B	N
ATOM	7672 CB GLU	237	81. 657 60. 643 , 26. 041 1. 00 15. 93 B	C
ATOM	7673 CG GLU	237	82. 307 60. 993 24. 708 1. 00 20. 06 B	C . C
ATOM	7674 CD GLU	237	81. 311 61. 541 23. 682 1. 00 24. 67 B	C
ATOM	7675 OE1 GLU	237	80. 133 61. 125 23. 713 1. 00 27. 11 B	0
ATOM	7676 OE2 GLU	237	81. 706 62. 377 22. 832 1. 00 25. 71 B	0
ATOM	7677 C GLU	237	81. 902 59. 898 28. 407 1. 00 12. 26 B	Č
ATOM	7.678 O GLU	237	81. 473 58. 759 28. 569 1. 00 12. 02 B	ŏ
ATOM ATOM	7679 N TYR	238	81. 768 60. 860 29. 310 1. 00 12. 67 B	Ň
ATOM	7680 CA TYR 7681 CB TYR	238	81. 044 60. 630 30. 550 1. 00 13. 08 B	Ĉ
ATOM		238	81. 903 59. 816 31. 534 1. 00 11. 88 B	C
ATOM	7682 CG TYR 7683 CD1 TYR	238	83. 201 60. 458 31. 954 1. 00 15. 20 B	C
ATOM	7684 CE1 TYR	238	83. 250 61. 347 33. 026 1. 00 15. 46 B	C
ATOM	7685 CD2 TYR	238	84. 458 61. 920 33. 430 1. 00 15. 78 B	C .
ATOM	7686 CE2 TYR	238 238	84. 390 60. 160 31. 291 1. 00 14. 07 B	C ·
ATOM	7687 CZ TYR	238	85. 592 60. 727 31. 683 1. 00 14. 24 B	C C C
ATOM	7688 OH TYR	238	85. 623 61. 606 32. 751 1. 00 13. 94 B	C
ATOM	7689 C TYR	238	86. 818 62. 173 33. 129 1. 00 12. 45 B 80. 583 61. 944 31. 163 1. 00 13. 53 B	0
ATOM	7690 O TYR	238	01 005 00 000	C
ATOM	7691 N SER	239	70 F00 64 60F	0
ATOM	7692 CA SER	239	70 040 00 045	N
			79.040 63.047 32.684 1.00 13.89 B UBSTITUTE SHEET (RULE 26)	С

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			FIG. 4	- 158	(Continued)
ATOM 77	7693 CB SER 7694 OG SER 7695 C SER 7696 O SER 7697 N PHE 7698 CA PHE 7699 CB PHE 7700 CG PHE 7701 CD1 PHE 7702 CD2 PHE 7704 CE2 PHE 7705 CZ PHE 7706 C PHE 7707 O PHE 7707 CB TYR 7710 CB TYR 7710 CB TYR 7711 CG TYR 7712 CD1 TYR 7712 CD1 TYR 7713 CE1 TYR 7714 CD2 TYR 7715 CE2 TYR 7716 CZ TYR 7717 OH TYR 7718 C SER 7720 N SER 7721 CA SER 7720 N SER 7721 CA SER 7721 CA SER 7722 CB SER 7723 OG SER 7724 C SER 7725 O SER 7726 N ASP 7727 CA ASP 7728 CB ASP 7729 CG ASP 7720 N SER 7721 CA SER 7721 CA SER 7722 CB SER 7723 OG SER 7724 C SER 7725 O SER 7726 N ASP 7727 CA ASP 7728 CB ASP 7729 CG ASP 7720 N SER 7721 CA SER 7723 OG SER 7724 C SER 7725 O SER 7726 N ASP 7727 CA ASP 7728 CB ASP 7729 CG ASP 7730 OD1 ASP 7731 OD2 ASP 7731 OD2 ASP 7732 CG GLU 7733 CG GLU 7735 CA GLU 7736 CB GLU 7737 CG GLU 7738 CD GLU 7737 CG GLU 7738 CD GLU 7739 OE1 GLU 7739 OE1 GLU 7730 OE2 GLU 7730 OE2 GLU 7731 OE2 GLU 7731 CG GLU 7731 CG GLU 7732 CG GLU 7733 CG GLU 7734 CG GLU 7735 CA GLU 7736 CB GLU 7737 CG GLU 7738 CD GLU 7739 OE1 GLU 7739 OE1 GLU 7730 OE2 GLU 7731 CG GLU	239 239 239 239 239 240 240 240 240 240 240 240 241 241 241 241 241 241 241 241 241 241	77. 597 62. 783 76. 800 62. 496 79. 775 63. 547 80. 361 62. 775 79. 737 64. 860 80. 313 65. 493 81. 543 66. 325 82. 422 66. 591 83. 325 65. 629 82. 312 67. 781 84. 108 65. 846 83. 087 68. 009 83. 988 67. 039 79. 184 66. 403 78. 671 67. 232 78. 785 66. 231 77. 683 67. 002 76. 912 66. 125 76. 480 64. 848 75. 393 64. 832 75. 051 63. 678 77. 215 63. 674 76. 883 62. 512 75. 801 62. 523 75. 489 61. 395 78. 100 68. 299 77. 311 69. 23	33. 085	C O C O N C C C C C C C C C C C C C C C

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				٠.	ĖI	~ 1 -	160			(Continued)
					L I (G. 4 -	100			
ATOM	7791	NZ	LYS	250	84. 262	70. 465	31.442	1.00 26.19	D	N
ATOM	7792	C	LYS	250 250	79. 215				B B	N . C
ATOM	7793	0	LYS	250 250		67.313	31.040	1.00 17.64 1.00 20.20	В	
ATOM	7794	N	THR		79.348	66. 409	31.867			0 N
ATOM	7795	CA	THR	251 251	79. 478 79. 978	67. 160 65. 905	29. 750 29. 234	1.00 15.06	B B	
ATOM	7796	CB	THR	251 251		65. 537	27.896	1.00 14.91		C C
ATOM	7797	0G1	THR		79. 317 77. 965	65. 144		1.00 13.86	B B	
ATOM	7798		THR	251 251	80. 058	64. 389	28. 128 27. 227	1.00 14.97 1.00 13.23	В	C C
ATOM	7799	CGZ	THR	251	81.473	66.016		1.00 15.25	В	C
ATOM	7800		THR			66.831	29. 015 28. 227		В	0
ATOM	7801	O N	VAL	251 252	81.934 82.231	65. 194		1.00 18.88	В	N N
ATOM	7802	CA	VAL	252 252	83. 675	65. 195	29.720	1.00 15.28 1.00 15.13	В	
ATOM	7803	CB	VAL	252	84. 335	64.717	29. 578 30. 882	1.00 13.13	В	C
ATOM	7804	CG1		252	85. 827	64. 580	30. 706	1.00 13.04	В	C
ATOM	7805	CG2		252	84. 012	65. 701	31.991	1.00 10.22	В	C C
ATOM	7806	C	VAL	252	84. 027	64. 264	28. 422	1.00 11.83	В	C
ATOM	7807	0	VAL	252	83. 472	63. 173	28. 304	1.00 17.21	В	0
ATOM	7808	N	ARG	253	84. 929	64.710	27. 557	1.00 17.34	. В	N N
ATOM	7809	CA	ARG	253 253	85. 349	63.922	26. 403	1.00 20.46	В	Č
ATOM	7810	CB	ARG	253 253	84. 822	64.560	25. 113	1.00 20.40	В	C
ATOM	7811	CG	ARG	253	83. 399	64. 137	24. 755	1.00 26.72	В	C
ATOM	7812	CD	ARG	253	82. 847	64. 920	23. 578	1.00 28.87	В	Č
ATOM	7813	NE	ARG	253	82. 176	66. 132	24. 033	1.00 36.20	В	N
ATOM	7814	CZ	ARG	253	80. 870	66. 221	24. 278	1.00 38.47	В	C
ATOM	7815		ARG	253	80. 084	65. 164	24. 099	1.00 39.84	В	N
ATOM	7816		ARG	253	80. 352	67.360	24. 727	1.00 37.97	В	N ·
ATOM	7817	C	ARG	253	86. 863	63.863	26. 389	1.00 19.71	В	Č
ATOM	7818	ŏ	ARG	253	87. 520	64.886	26. 246	1.00 21.87	B.	ŏ
ATOM	7819	Ň	VAL	254	87. 404	62.656	26. 538	1.00 18.34	В	N
ATOM	7820	ĊA	VAL	254	88. 847	62.434	26. 594	1.00 15.15	B	Č
ATOM	7821	CB	VAL	254	89. 257	61.924	27. 994	1.00 16.16	В	č
ATOM	7822		VAL	254	90. 771	61.759	28. 081	1.00 15.18	В	č
ATOM	7823		VAL	254	88. 736	62.868	29.065	1.00 16.46	B	Č
ATOM	7824	C	VAL	254	89. 313	61.397	25. 585	1.00 14.67	В	Č
· ATOM	7825	0	VAL	254	88. 806	60. 272	25. 566	1.00 14.87	В	. 0
ATOM	7826	Ň	PRO	255	90. 281	61.757	24. 726	1.00 13.62	B	N
ATOM	7827	CD	PR0	255	90.872	63.081	24. 472	1.00 12.90	B	Č
ATOM	7828	CA	PRO	255	90.760	60.777	23. 746	1.00 12.62	B	Č ·
ATOM	7829	CB	PR ₀	255	91.786	61.566	22. 933	1.00 11.40	B	č
ATOM	7830	CG	PRO	255	91. 263	62.969	23.013	1.00 11.65	B	č
ATOM	7831	C	PR0	255	91.379	59.645	24. 553	1.00 12.46	B	č
ATOM	7832	0	PRO	255	92. 355	59.831	25. 282	1.00 13.25	B	ŏ
ATOM	7833	N	TYR	256	90. 796	58.469	24. 414	1.00 12.53	B	Ň
ATOM	7834	CA	TYR	256	91.217	57.306	25.161	1.00 12.05	B	Ĉ
ATOM	7835	CB	TYR	256	90. 319	57. 205	26. 398	1.00 12.42	B	č
ATOM	7836	CG	TYR	256	90.608	56.082	27.360	1.00 14.53	B	Č
ATOM	7837	CD1	TYR	256	91.021	56.355	28.662	1.00 16.44	B	Č
ATOM	7838	CE1	TYR	256	91. 192	55.337	29.596	1.00 17.38	В	Č
ATOM	7839	CD2	TYR	256	90. 382	54.752	27.010	1.00 15.31	В	C

		— — — —	(Continued)
		FIG. 4-161	(33-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-
ATOM 7841 CZ ATOM 7842 OH ATOM 7843 C ATOM 7844 O ATOM 7845 N ATOM 7846 CD ATOM 7847 CA ATOM 7849 CG ATOM 7850 C ATOM 7851 O ATOM 7852 N ATOM 7852 N ATOM 7853 CA ATOM 7854 CB ATOM 7855 CG ATOM 7856 CD ATOM 7857 CE ATOM 7858 NZ ATOM 7860 O ATOM 7860 O ATOM 7861 N ATOM 7862 CA ATOM 7863 CB ATOM 7863 CB ATOM 7863 CB ATOM 7864 C	2 TYR 256 PRO 257 PRO 258 LYS 258	90. 949 54. 030 29. 232 1. 00 16. 54 91. 068 53. 042 30. 176 1. 00 17. 03 91. 040 56. 094 24. 263 1. 00 11. 63 89. 923 55. 765 23. 870 1. 00 13. 76 92. 141 55. 415 23. 924 1. 00 10. 78 93. 535 55. 786 24. 231 1. 00 9. 21 92. 098 54. 229 23. 068 1. 00 9. 97 93. 473 54. 233 22. 438 1. 00 8. 95 94. 326 54. 657 23. 606 1. 00 8. 91 91. 859 52. 949 23. 869 1. 00 11. 12 92. 694 52. 556 24. 681 1. 00 9. 90 90. 723 52. 300 23. 648 1. 00 11. 97 90. 444 51. 057 24. 353 1. 00 13. 52 88. 930 50. 855 24. 492 1. 00 15. 66 88. 305 51. 808 25. 522 1. 00 14. 41 86. 801 51. 730 25. 552 1. 00 14. 41 86. 801 51. 730 25. 552 1. 00 14. 62 91. 101 49. 934 23. 571 1. 00 16. 07 91. 227 48. 760 24. 178 1. 00 16. 07 91. 227 48. 760 24. 178 1. 00 16. 07 91. 227 48. 760 24. 178 1. 00 16. 07 91. 227 48. 760 24. 178 1. 00 16. 09 91. 476 47. 476 22. 045 1. 00 16. 09	3
ATOM 7864 C ATOM 7865 O ATOM 7866 N ATOM 7867 CA ATOM 7868 C ATOM 7869 O ATOM 7870 N ATOM 7871 CA ATOM 7872 CB ATOM 7873 C ATOM 7874 O ATOM 7875 N ATOM 7876 CA ATOM 7877 CB ATOM 7877 CB ATOM 7878 CG1 ATOM 7879 CG2 ATOM 7880 C	ALA 259 ALA 259 ALA 259 GLY 260 GLY 260 GLY 260 GLY 261 ALA 261 ALA 261 ALA 261 ALA 261 ALA 261 ALA 262 VAL 262 VAL 262 VAL 262 VAL 262 VAL 262 VAL 262	91. 564 46. 356 24. 261 1. 00 14. 32 B 91. 476 47. 476 22. 045 1. 00 16. 09 B 90. 293 47. 415 21. 710 1. 00 15. 64 B 92. 477 47. 428 21. 172 1. 00 15. 95 B 92. 221 47. 269 19. 754 1. 00 15. 99 B 91. 841 48. 523 18. 982 1. 00 17. 08 B 91. 781 48. 488 17. 752 1. 00 18. 87 B 91. 587 49. 629 19. 673 1. 00 14. 62 B 91. 198 50. 851 18. 983 1. 00 14. 89 B 90. 557 51. 830 19. 963 1. 00 13. 58 B 92. 379 51. 509 18. 292 1. 00 17. 12 B 93. 489 50. 986 18. 298 1. 00 20. 05 B 92. 135 52. 662 17. 686 1. 00 17. 34 B 93. 192 53. 384 17. 004 1. 00 16. 00 B 92. 614 54. 371 15. 947 1. 00 14. 51 B 93. 717 55. 252 15. 383 1. 00 13. 59 B 91. 970 53. 596 14. 820 1. 00 10. 82 B 93. 984 54. 150 18. 055 1. 00 17. 31 B	C C O N C C C C C C C C C C C C C C C C
ATOM 7882 N A ATOM 7883 CA A ATOM 7884 CB A ATOM 7885 CG A ATOM 7886 OD1 A ATOM 7887 ND2 A		93. 432 54. 973 18. 786 1. 00 20. 51 B 95. 275 53. 856 18. 128 1. 00 16. 87 B 96. 190 54. 493 19. 068 1. 00 17. 45 B 97. 406 53. 595 19. 292 1. 00 17. 58 B 97. 230 52. 629 20. 437 1. 00 20. 08 B 97. 919 51. 606 20. 500 1. 00 19. 88 B 96. 329 52. 950 21. 365 1. 00 18. 44 B 96. 706 55. 827 18. 533 1. 00 18. 01 B	O N C C C O N C

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						(Continued)
					FIG. 4-162	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7889 7890 7891 7892 7893 7894 7895 7896 7897 7898 7899 7900		ASN PRO PRO PRO PRO PRO PRO THR THR THR	263 264 264 264 264 264 265 265 265 265	96. 578 56. 134 17. 345 1. 00 19. 39 B 97. 288 56. 646 19. 413 1. 00 17. 06 B 97. 357 56. 546 20. 883 1. 00 15. 68 B 97. 819 57. 926 18. 950 1. 00 15. 10 B 98. 089 58. 676 20. 251 1. 00 14. 78 B 98. 411 57. 569 21. 214 1. 00 14. 94 B 99. 105 57. 605 18. 198 1. 00 15. 50 B 99. 669 56. 527 18. 369 1. 00 15. 27 B 99. 560 58. 521 17. 354 1. 00 16. 21 B 100. 796 58. 305 16. 617 1. 00 15. 30 B 100. 647 58. 677 15. 132 1. 00 15. 20 B 100. 081 59. 983 15. 029 1. 00 17. 05 B	0 N C C C C C O N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7901 7902 7903 7904 7905 7906 7907 7908 7909	C O N CA CB CG1	THR THR VAL VAL VAL VAL VAL VAL VAL	265 265 265 266 266 266 266 266	99. 747 57. 687 14. 415 1. 00 10. 60 B 101. 818 59. 211 17. 279 1. 00 16. 13 B 101. 454 60. 126 18. 007 1. 00 16. 83 B 103. 095 58. 971 17. 030 1. 00 17. 64 B 104. 118 59. 781 17. 667 1. 00 17. 49 B 104. 626 59. 060 18. 930 1. 00 15. 28 B 105. 224 57. 714 18. 538 1. 00 12. 10 B 105. 642 59. 921 19. 666 1. 00 12. 62 B	C C O N C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7910 7911 7912 7913 7914 7915 7916	O N CA CB CG CD CE	VAL LYS LYS LYS LYS LYS LYS	266 267 267 267 267 267 267	105. 312 60. 112 16. 769 1. 00 19. 23 B 105. 693 59. 331 15. 893 1. 00 18. 24 B 105. 889 61. 287 17. 003 1. 00 20. 19 B 107. 058 61. 756 16. 272 1. 00 19. 42 B 106. 678 62. 855 15. 291 1. 00 19. 76 B 105. 786 62. 413 14. 168 1. 00 21. 59 B 105. 452 63. 605 13. 291 1. 00 23. 15 B 104. 593 63. 205 12. 119 1. 00 23. 47 B	C O N C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7917 7918 7919 7920 7921 7922 7923 7924 7925	NZ C O N CA CB CG CD1 CD2		267 267 267 268 268 268 268 268 268	104. 225 64. 402 11. 334 1. 00 27. 20 B 108. 032 62. 334 17. 288 1. 00 19. 59 B 107. 618 62. 826 18. 336 1. 00 20. 86 B 109. 322 62. 275 16. 984 1. 00 19. 32 B 110. 325 62. 818 17. 882 1. 00 18. 94 B 111. 350 61. 757 18. 259 1. 00 17. 47 B 112. 186 62. 131 19. 444 1. 00 16. 21 B 111. 601 62. 290 20. 692 1. 00 16. 98 B	N C O N C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7926 7927 7928 7929 7930 7931 7932 7933 7934 7935 7936 7937	CE1 CE2 CZ C O N CA CB	PHE	268 268 268 268 269 269 269 269 269 269 269	113.555 62.327 19.313 1.00 16.35 B 112.368 62.639 21.797 1.00 18.80 B 114.332 62.674 20.405 1.00 17.68 B 113.737 62.832 21.655 1.00 18.66 B 111.016 63.979 17.192 1.00 20.34 B 111.114 64.016 15.968 1.00 21.73 B 111.491 64.931 17.981 1.00 20.76 B 112.152 66.105 17.435 1.00 20.74 B 111.141 67.239 17.222 1.00 19.80 B 110.070 66.937 16.216 1.00 21.88 B 110.332 67.019 14.853 1.00 22.75 B 108.785 66.605 16.631 1.00 23.20 B 109.326 66.781 13.912 1.00 21.98 B	C C C C C C C C C C C C C C C C C C C

•			FIG. 4-163	(Continued)
4.MO)			•	
ATOM ATOM			107. 771 66. 364 15. 700 1. 00 23. 06	ВС
ATOM			108. 044 66. 454 14. 337 1. 00 22. 44	B C
ATOM			113. 209 66. 606 18. 402 1. 00 21. 66 113. 127 66. 376 19. 613 1. 00 21. 27	B C
ATOM				B 0
ATOM	7943 CA VAL		114. 195 67. 305 17. 858 1. 00 21. 99 115. 239 67. 896 18. 667 1. 00 23. 26	B N
ATOM	7944 CB VAL		116. 527 67. 062 18. 635 1. 00 23. 10	B C B C
ATOM			117.517 67.624 19.630 1.00 23.57	B C
ATOM			116. 219 65. 609 18. 985 1. 00 23. 02	B Č
MOTA	7947 C VAL		115. 495 69. 285 18. 095 1. 00 25. 32	B C
ATOM ATOM	7948 0 VAL		115. 600 69. 460 16. 880 1. 00 26. 00	ВО
ATOM	7949 N VAL 7950 CA VAL		115.561 70.278 18.973 1.00 26.96	B N
ATOM	7950 CA VAL 7951 CB VAL		115. 794 71. 650 18. 546 1. 00 27. 45	B C
ATOM	7952 CG1 VAL		114.516 72.514 18.714 1.00 28.95 114.096 72.563 20.177 1.00 28.40	B C
ATOM	7953 CG2 VAL		111 500 50.10	B C
ATOM	7954 C VAL	271	114. 769 73. 915 18. 186 1. 00 29. 54 116. 926 72. 258 19. 363 1. 00 27. 39	B C
ATOM	7955 O VAL	271	117. 094 71. 935 20. 536 1. 00 26. 71	B C B O
ATOM	7956 N ASN	272	117. 706 73. 128 18. 728 1. 00 27. 87	B N
ATOM	7957 CA ASN	272	118. 828 73. 788 19. 383 1. 00 27. 39	B C
ATOM	7958 CB ASN	272	119. 951 74. 056 18. 378 1. 00 27. 64	B C
ATOM ATOM	7959 CG ASN	272	121. 179 74. 672 19. 031 1. 00 29. 11	B Č
ATOM	7960 OD1 ASN 7961 ND2 ASN	272	121. 094 75. 706 19. 696 1. 00 28. 35	ВО
ATOM	7962 C ASN	$\begin{array}{c} 272 \\ 272 \end{array}$	122. 330 74. 036 18. 841 1. 00 30. 38	B N
ATOM	7963 0 ASN	272	118. 347 75. 104 19. 972 1. 00 27. 43	ВС
ATOM	7964 N THR	273	117. 943 76. 012 19. 243 1. 00 27. 41 118. 397 75. 208 21. 292 1. 00 27. 62	B 0
ATOM	7965 CA THR	273		B N
ATOM	7966 CB THR	273	117. 938 76. 411 21. 959 1. 00 28. 77 117. 509 76. 100 23. 400 1. 00 27. 46	B C
ATOM	7967 OG1 THR	273	118. 653 75. 727 24. 181 1. 00 28. 06	B C B O
ATOM	7968 CG2 THR	273	116.510 74.960 23.403 1.00 26.33	B C
ATOM	7969 C THR	273	118. 988 77. 516 21. 967 1. 00 31. 23	B C
ATOM ATOM	7970 O THR	273	118.669 78.680 22.208 1.00 32.66	B 0
ATOM	7971 N ASP 7972 CA ASP	274	120. 239 77. 157 21. 698 1. 00 32. 45	B N
ATOM	7973 CB ASP	274 274	121.315 78.139 21.676 1.00 33.79	B C
ATOM	7974 CG ASP	274	122. 671 77. 446 21. 775 1. 00 34. 63 123. 019 77. 049 23. 193 1. 00 36. 82	B C
ATOM	7975 OD1 ASP	274	104 045 80 000	B C
ATOM	7976 OD2 ASP	274	100 005 55 100	B 0
ATOM	7977 C ASP	274	122. 267 77. 430 24. 117 1. 00 37. 18 121. 277 78. 996 20. 419 1. 00 35. 09	B 0
ATOM	7978 O ASP	274	121. 899 80. 058 20. 366 1. 00 34. 53	B C B O
ATOM	7979 N SER	275	120. 540 78. 542 19. 412 1, 00 35, 53	B N
ATOM	7980 CA SER	275	120. 456 79. 279 18. 166 1. 00 37. 39	B C
ATOM	7981 CB SER	275	121. 096 78. 462 17. 051 1. 00 36. 90	B C
ATOM ATOM	7982 OG SER 7983 C SER	275	120. 476 77. 197 16. 948 1. 00 41. 05	B 0
ATOM	7983 C SER 7984 O SER	275 275	119. 030 79. 652 17. 781 1. 00 39. 58	B C
ATOM	7985 N LEU	275 276	118. 580 79. 355 16. 673 1. 00 40. 66	B 0
ATOM	7986 CA LEU	276	118. 323 80. 311 18. 695 1. 00 40. 56 116. 949 80. 732 18. 443 1. 00 40. 53	B N
	7. 200		116. 949 80. 732 18. 443 1. 00 40. 53	B C

			a	(Continue	d)
			FIG. 4-164		
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	7987 CB LET 7988 CG LET 7989 CD1 LET 7990 CD2 LET 7991 C LET 7992 O LET 7993 N SEF 7994 CA SEF 7995 CB SEF 7996 OG SEF 7997 C SEF 7998 O SEF 7999 N SEF 8000 CA SEF 8001 CB SEF 8002 OG SEF 8003 C SEF 8004 O SEF 8005 N VAL 8006 CA VAL 8007 CB VAL	7 276 7 276 7 276 8 277 8 277 8 277 9 277 9 277 9 277 9 278 9 278	116.076 80.425 19.664 1.00 38.58 116.002 78.958 20.097 1.00 36.34 115.319 78.876 21.445 1.00 35.16 115.261 78.134 19.057 1.00 32.57 116.914 82.229 18.140 1.00 41.99 117.675 83.002 18.721 1.00 41.16 116.029 82.634 17.233 1.00 44.02 115.916 84.044 16.863 1.00 46.53 116.489 84.277 15.462 1.00 48.49 116.268 85.618 15.044 1.00 50.90 114.494 84.586 16.902 1.00 46.23 113.529 83.856 16.701 1.00 46.82 114.378 85.884 17.148 1.00 49.4 113.081 86.535 17.202 1.00 47.82 113.617 87.759 19.234 1.00 48.09 113.617 87.759 19.234 1.00 48.26	B C B C B C B C B C B C B C B C B C B C	
ATOM ATOM	8007 CB VAL 8008 CG1 VAL 8009 CG2 VAL	279 279 279	113. 709 87. 454 11. 089 1. 00 50. 61	B C B C	
ATOM	8010 C VAL	279	114. 902 88. 464 13. 037 1. 00 50. 28 112. 340 85. 606 12. 941 1. 00 48. 52	B C B C	
ATOM ATOM	8011 O VAL 8012 N THR	279 280	111. 130 85. 433 13. 082 1. 00 49. 49 113. 145 84. 708 12. 380 1. 00 47. 70	B O B N	
ATOM ATOM	8013 CA THR 8014 CB THR	280	112. 651 83. 432 11. 872 1. 00 46. 64	B C	
ATOM	8015 OG1 THR	280 280	113. 719 82. 709 11. 032 1. 00 47. 86 113. 179 81. 479 10. 531 1. 00 48. 07	B C B O	
ATOM	8016 CG2 THR	280	114. 946 82. 399 11. 883 1. 00 47. 49	B C	
ATOM ATOM	8017 C THR 8018 O THR	280 280	112. 238 82. 484 12. 992 1. 00 45. 40	B C	
ATOM	8019 N ASN	281	112. 586 82. 677 14. 155 1. 00 44. 24 111. 499 81. 447 12. 622 1. 00 45. 09	B O B N	
ATOM	8020 CA ASN	281	111.040 80.454 13.581 1.00 44.81	B C	
ATOM	8021 CB ASN	281	109. 744 79. 815 13. 089 1. 00 46. 08	B C	
ATOM ATOM	8022 CG ASN 8023 OD1 ASN	281 281	108. 592 80. 786 13. 096 1. 00 48. 90 108. 351 81. 455 14. 101 1. 00 49. 62	B C	
ATOM	8024 ND2 ASN	281	108. 351 81. 455 14. 101 1. 00 49. 62 107. 873 80. 868 11. 984 1. 00 52. 14	B O B N	
ATOM	8025 C ASN	281	112.088 79.379 13.812 1.00 43.47	B N B C	
ATOM	8026 O ASN	281	112.874 79.065 12.919 1.00 44.44	B 0	
ATOM ATOM	8027 N ALA 8028 CA ALA	282 282	112. 100 78. 823 15. 019 1. 00 41. 76	B N	
ATOM	8029 CB ALA	282	113. 045 77. 773 15. 371 1. 00 38. 62 112. 795 77. 301 16. 792 1. 00 37. 75	B C B C	
ATOM	8030 C ALA	282	112. 863 76. 619 14. 403 1. 00 37. 34	B C B C	
ATOM	8031 O ALA	282	111.797 76.463 13.815 1.00 36.86	B 0	
ATOM ATOM	8032 N THR	283	113. 905 75. 816 14. 231 1. 00 36. 50	B N	
ATOM	8033 CA THR 8034 CB THR	283 283	113. 828 74. 672 13. 335 1. 00 35. 84 114. 867 74. 772 12. 218 1. 00 37 70	B C	
ATOM	8035 OG1 THR	283	114.867 74.772 12.218 1.00 37.70 114.665 75.994 11.495 1.00 41.71	B C B O	
				·	

						C 4	1.0	_		(Continued)
					P I	G. 4	- 10;)		
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8060 8061 8062 8063 8064 8065 8066 8067 8068 8069 8070 8071 8072 8073 8074	C O N CA CB CG CG CD N CA CB CG CG CD N CA CB CG CG CD N CA CB CG CG CC	THE SERRE ELECTION OF STATE OF	283 283 284 284 284 284 285 285 285 285 285 285 286 286 286 286 286 286 286 287 287 287 287 287 287 287 287 287 287	114. 736 114. 074	73. 403 73. 263 72. 482 71. 230 70. 893 71. 761 70. 090 69. 865 69. 367 68. 241 68. 037 66. 979 69. 350	11. 268 14. 128 14. 77 14. 073 14. 800 15. 507 16. 608 13. 883 12. 850 14. 260 13. 457 13. 546 12. 548 13. 250 14. 428 13. 976 15. 168	5 1.00 37.32 1.00 33.58 1.00 34.31 3 1.00 32.05 1.00 30.43 1.00 28.61 1.00 29.31 1.00 30.34 1.00 31.22 1.00 29.19 1.00 29.80 1.00 29.35 1.00 29.12 1.00 30.38 1.00 34.47 1.00 28.14 1.00 30.23	B B B B B B B B B B B B B B B B B B B	C C O N C C C C C C C O N C C C C C C O N C C C C
ATOM	8073	0G1	THR	288	113. 172	58.516	10.539	1.00 26.37	В	0
ATOM	8075	C	THR	288	112.529	57.741	10. 593 13. 335	1.00 25.25 1.00 26.85	B B	C C
ATOM ATOM			THR ALA	288 289	113.687 .111.484	57. 379 57. 011	13. 503 13. 702	1.00 27.04 1.00 28.37	B B	0
ATOM ATOM	8078	CA	ALA ALA	289 289	111.638	55.705	14.325	1.00 27.90	В	N C
ATOM	8080	C	ALA	289	110. 271 112. 348	55. 151 54. 740	14. 710 13. 380	1.00 26.91 1.00 27.44	В	C C
ATOM ATOM			ALA PRO	289 290	112. 550 112. 758	55. 038 53. 577	12. 205 13. 895	1.00 28.30 1.00 26.01	B B	0
ATOM ATOM		CD :	PRO PRO	290 290	112. 903 113. 445	53. 280 52. 569	15. 328 13. 089	1.00 24.74	B B	N C
					SUBSTITUTE			1.00 25.29 (ם	С

				94. 194.						
					TO T (~ 4	166			(Continued)
					ГІ	G. 4-	100			
ATOM	8085	CB	PRO	290	113. 949	51.587	14. 138	1.00 25.76	В	С
ATOM	8086	CG	PRO	290	114. 151	52.467	15.342		В	č
ATOM	8087	Č	PRO	290	112. 465	51.931	12.110		В	č
ATOM	8088	Ō	PRO	290	111. 255		12. 330		B	ŏ
ATOM	8089	N	ALA	291	112. 988	51.345	11.038		B	Ň
ATOM	8090	CA	ALA	291	112.143	50.730	10.024		B	Ċ
ATOM	8091	CB	ALA	291	112.987	50. 271	8.846	1.00 26.28	В	Č
ATOM	8092	C	ALA	291	111. 337	49.568	10.573	1.00 27.18	В	С
ATOM	8093	0	ALA	291	110. 203	49.331	10.145	1.00 27.46	В	0
ATOM	8094	N	SER	292	111.916	48.843	11.521	1.00 27.54	В	N
ATOM	8095	CA	SER	292	111.220		12.103	1.00 28.19	В	C
ATOM	8096	CB	SER	292	112.161	46.892	12.993	1.00 28.00	В	C
ATOM	8097	0G	SER	292	112. 525	47.626	14. 145	1.00 32.22	В	0
ATOM	8098	C	SER	292	110.027	48. 182	12.922	1.00 28.13	В	C
ATOM	8099	0	SER	292	109.176	47. 376	13. 307	1.00 29.52	В	0
MOTA	8100	N	MET	293	109.976	49. 487	13. 190	1.00 25.00	В	N
ATOM	8101	CA	MET	293	108. 881	50.072	13. 955	1.00 24.80	В	C
ATOM ATOM	8102	CB CG	MET	293	109. 387	51.173	14.892	1.00 24.61	В	C
ATOM	8103 8104	SD	MET MET	293 293	110. 231	50. 703	16.060	1.00 26.88	В	C
ATOM	8105	CE	MET	293 293	109. 323 110. 457	49. 647 48. 319	17. 189	1.00 27.80	В	S
ATOM	8106	C	MET	293	107. 836	50. 677	17. 438 13. 027	1.00 25.74 1.00 24.57	В	C
ATOM	8107	0	MET	293	106.641	50. 528	13. 027	1.00 24.37	B B	C 0
ATOM	8108	Ň	LEU	294	108. 292	51. 360	11.983	1.00 23.32	В	N N
ATOM	8109	ĊA	LEU	294	107. 393	52.008	11.041	1. 00 23. 80	В	C
ATOM	8110	CB	LEU	294	108. 183	52. 930	10.114	1. 00 23. 40	В	č
ATOM	8111	CG	LEU	294	108. 945	54.072	10.786	1.00 24.87	B	č
ATOM	8112	CD1	LEU	294	109.806	54. 787	9.758	1.00 22.08	B	č
ATOM	8113	CD2	LEU	294	107.958	55.037	11.440	1.00 23.08	B	Č
ATOM	8114	C	LEU	294	106.540	51.059	10.204	1.00 23.95	В	Č
ATOM	8115	0	LEU	294	105.714	51.510	9.422	1.00 25.36	В	0
ATOM	8116	N	ILE	295	106.724	49. 754	10.357	1.00 23.92	В	N
ATOM	8117	CA	ILE	295	105.923	48.812	9. 580	1.00 25.26	В	C
ATOM	8118	CB	ILE	295	106.601	47.444	9.453	1.00 26.06	В	С
ATOM	8119		ILE	295	107. 972	47. 595	8.812	1.00 26.54	В	C
ATOM	8120		ILE	295	106.698	46. 796	10.831	1.00 24.44	В	C
ATOM	8121		ILE	295	107. 211	45. 388	10.789	1.00 28.37	В	C
ATOM	8122	C	ILE	295	104.564	48. 575	10. 221	1.00 26.01	В.	C
ATOM ATOM	8123 8124	O N	ILE GLY	295	103. 805	47. 712	9.775	1.00 28.75	В	0
ATOM	8125	CA	GLY	296	104. 263	49. 328		1.00 24.77	В	N
ATOM	8126	C	GLY	296 296	102.992	49. 167 50. 040	11.951	1.00 22.28	В	C
ATOM	8127	Ö	GLY	296	102. 908 103. 820	50. 818	13. 182 13. 447	1. 00 21. 29 1. 00 20. 80	В	C
ATOM	8128	N	ASP	297	103.820	49. 920	13. 447	1.00 20.80	B B	0
ATOM	8129	CA	ASP	297	101.654	50. 718	15. 141	1.00 20.38	В	N C
ATOM	8130	CB	ASP	297	100.366	50. 339	15. 874	1.00 20.14	В	C C
ATOM	8131	CG	ASP	297	99. 109	50. 665	15.078	1.00 22.60	В	C
ATOM	8132	0D1		297	98. 016	50. 234	15.502	1.00 25.00	В	0
ATOM	8133	0D2		297	99. 200	51.350	14.041	1.00 22.18	В	0 .
									_	-

	٠				FIG. 4-167	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8134 8135 8136 8137 8138 8140 8141 8142 8143 8144 8145 8146 8150 8151 8152 8153 8154 8155 8156 8157 8158 8159 8160 8161 8162 8163 8164 8165 8166 8167 8168	O N CA CB CG CD1 CC CZ CZ OH C C CD1 CC CD1 CC	HIS HIS HIS HIS HIS HIS HIS HIS HIS TYR TYR TYR TYR	297 297 298 298 298 298 298 298 298 299 299 299	102. 845 50. 481 16. 065 1. 00 20. 31 B 103. 419 49. 390 16. 096 1. 00 20. 82 B 103. 220 51. 508 16. 814 1. 00 16. 87 B 104. 335 51. 384 17. 734 1. 00 16. 48 B 105. 669 51. 399 16. 968 1. 00 14. 91 B 105. 868 52. 628 16. 137 1. 00 12. 24 B 106. 539 53. 775 16. 391 1. 00 10. 39 B 105. 264 52. 802 14. 909 1. 00 11. 35 B 105. 551 54. 005 14. 445 1. 00 11. 25 B 106. 323 54. 616 15. 326 1. 00 11. 96 B 104. 274 52. 560 18. 693 1. 00 15. 50 B 105. 127 52. 539 19. 706 1. 00 15. 50 B 105. 163 53. 599 20. 698 1. 00 15. 50 B 105. 163 53. 599 22. 047 1. 00 14. 50 B 104. 64	CONCCCNCNCONCCCCCCCCONCCCCCONCC
ATOM ATOM ATOM	8169 8170 8171	SG C O	CYS CYS CYS	301 301 301	109. 922 51. 905 24. 722 1. 00 26. 11 B 109. 895 54. 842 26. 194 1. 00 20. 82 B 109. 816 54. 579 27. 395 1. 00 21. 62 B	C S C O
ATOM ATOM ATOM ATOM ATOM ATOM	8172 8173 8174 8175 8176 8177		ASP ASP ASP ASP ASP ASP	302 302 302 302 302 302	110.922 55.496 25.662 1.00 22.13 B 112.035 55.968 26.481 1.00 20.03 B 112.875 54.810 27.014 1.00 20.49 B 114.035 55.296 27.868 1.00 25.77 B 113.880 55.344 29.109 1.00 26.02 B 115.097 55.664 27.297 1.00 27.73 B	N C C C O O
ATOM ATOM ATOM ATOM ATOM	8178 8179 8180 8181 8182	C O N CA CB	ASP ASP VAL VAL VAL	302 302 303 303 303	112.959 56.894 25.711 1.00 20.08 B 113.367 56.596 24.586 1.00 19.30 B 113.302 58.010 26.343 1.00 20.41 B 114.188 59.000 25.756 1.00 20.36 B 113.435 60.316 25.470 1.00 19.97 B	C O . N C C

								•
				FIG. 4-1	168			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8183 8184 8185 8186 8187 8188 8190 8191 8192 8193 8194 8195 8199 8200 8201 8202 8203 8204 8205 8206 8207 8208 8209 8210	CG1 VAL CG2 VAL C VAL O VAL N THR CA THR CG1 THR CG2 THR CG2 THR CG3 TRP CG TRP CG TRP CCB TRP	303 303 303 304 304 304 304 305 305 305 305 305 305 305 305 305 305	114. 387 61. 347 112. 260 60. 043 115. 267 59. 251 114. 950 59. 568 116. 536 59. 112 117. 639 59. 313 118. 008 58. 002 116. 869 57. 496 119. 136 58. 242 118. 925 59. 851 119. 579 59. 159 119. 307 61. 069 120. 545 61. 643 120. 696 63. 114 119. 834 64. 751 118. 614 65. 413 120. 885 64. 928 118. 414 64. 232 117. 764 65. 077 118. 413 66. 242 120. 689 65. 746 119. 459 66. 395 121. 722 60. 875 121. 743 60. 552 122. 697 60. 591 123. 899 59. 864	24. 857 24. 540 26. 788 27. 939 26. 389 27. 332 28. 046 28. 751 29. 026 26. 729 25. 952 27. 102 26. 583 26. 975 26. 354 25. 150 24. 917 24. 243 25. 938 23. 812 23. 152 22. 943 27. 148 28. 338 26. 285 26. 673	1. 00 20. 23 1. 00 17. 52 1. 00 21. 02 1. 00 19. 39 1. 00 21. 38 1. 00 21. 48 1. 00 19. 77 1. 00 19. 55 1. 00 20. 57 1. 00 22. 96 1. 00 25. 30 1. 00 22. 41 1. 00 21. 86 1. 00 20. 21 1. 00 18. 79 1. 00 18. 79 1. 00 18. 79 1. 00 18. 65 1. 00 17. 49 1. 00 18. 37 1. 00 19. 16 1. 00 19. 59 1. 00 21. 43 1. 00 22. 21 1. 00 21. 63 1. 00 22. 53 1. 00 21. 31 1. 00 20. 65	B B B B B B B B B B B B B B B B B B B	C C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8211 8212 8213 8214 8215 8216 8217 8218 8220 8221 8222 8223 8224 8225 8226 8227 8228 8229 8230 8231	C ALA O ALA N THR CA THR CB THR OGI THR CG2 THR C THR O THR N GLN CA GLN CB GLN CB GLN CCB GLN	306 306 307 307 307 307 307 308 308 308 308 308 308 308 308 308 308	124. 975 60. 882 2 125. 675 60. 767 2 125. 086 61. 885 2 126. 057 62. 964 2 127. 285 62. 744 2 126. 894 62. 855 2 127. 892 61. 374 2 125. 397 64. 250 2 124. 177 64. 326 2 125. 699 66. 540 2 126. 762 67. 634 2 127. 301 67. 811 2 126. 256 68. 296 2 125. 116 68. 727 2 125. 126. 477 68. 290 2 125. 126. 66. 501 2 125. 370 65. 459 2 125. 370 65. 374 2	25. 533 27. 000 28. 007 26. 133 26. 284 25. 411 24. 040 25. 659 25. 731 25. 731 25. 479 25. 175 26. 574 27. 548 28. 754 27. 548 28. 754 29. 659 29.	1.00 20.65 1.00 21.97 1.00 20.32 1.00 23.85 1.00 24.42 1.00 25.33 1.00 19.34 1.00 25.73 1.00 26.09 1.00 24.49 1.00 22.95 1.00 21.20 1.00 23.08 1.00 23.08 1.00 21.02 1.00 25.59 1.00 26.16 1.00 25.99	B B B B B B B B B B B B B B B B B B B	C C O N C C C C O N C C C C C O N C C C C

		FIG. 4-169	(Continued)
ATOM 8233 CD CATOM 8234 OE1 CATOM 8235 OE2 CATOM 8236 C CATOM 8237 O CATOM 8238 N AATOM 8239 CA AATOM 8241 CG AATOM 8241 CG AATOM 8242 CD AATOM 8243 NE AATOM 8244 CZ AATOM 8245 NH1 AATOM 8246 NH2 AATOM 8247 C AATOM 8248 O AATOM 8249 N III ATOM 8250 CA III AT	309 ELU 309 ELU 309 ELU 309 ERG 310 ERG 311 ER 311 ER 311 ER 312 ER 313 ER 312 ER 313 ER 314 ER 315 ER 316	126. 925 67. 280 20. 774 1. 00 29. 27 128. 243 67. 637 20. 109 1. 00 31. 48 128. 614 66. 968 19. 115 1. 00 33. 35 128. 900 68. 593 20. 572 1. 00 32. 54 124. 939 63. 991 21. 004 1. 00 26. 83 124. 850 63. 712 19. 806 1. 00 28. 64 124. 674 63. 131 21. 982 1. 00 25. 93 124. 246 61. 765 21. 723 1. 00 24. 07 125. 357 60. 790 22. 121 1. 00 24. 28	B C C B C C C C C C C C C C C C C C C C
ATOM 8279 O GLM ATOM 8280 N TRP		111. 412 54. 045 22. 513 1. 00 22. 96 B 111. 984 52. 372 21. 108 1. 00 22. 35 B	O N

			•		. D. C. 4. 1.7.0	(Continued)
				-0.	FIG. 4-170	
ATOM ATOM ATOM ATOM ATOM ATOM	8281 8282 8283 8284 8285 8286	CE2 CE3	TRP TRP TRP TRP TRP TRP	315 315 315 315 315 315	110. 672 52. 262 20. 484 1. 00 21. 75 E 110. 769 52. 440 18. 968 1. 00 21. 09 E 111. 376 53. 741 18. 540 1. 00 21. 09 E 110. 678 54. 940 18. 176 1. 00 19. 81 E 111. 654 55. 901 17. 824 1. 00 20. 24 E 109. 325 55. 295 18. 113 1. 00 17. 16 E	3 C 3 C 3 C
ATOM ATOM ATOM ATOM ATOM ATOM	8287 8288 8289 8290 8291 8292	NE1 CZ2 CZ3	TRP TRP TRP TRP TRP TRP	315 315 315 315 315 315	112. 705 54. 018 18. 405 1. 00 21. 12 E 112. 880 55. 310 17. 974 1. 00 21. 84 E 111. 321 57. 197 17. 413 1. 00 18. 97 E 108. 992 56. 588 17. 704 1. 00 20. 13 E 109. 990 57. 522 17. 359 1. 00 19. 26 E 110. 118 50. 880 20. 790 1. 00 22. 37 E	B N B C B C
ATOM ATOM ATOM ATOM ATOM ATOM	8293 8294 8295 8296 8297 8298	O N CA CB CG CD1	TRP LEU LEU LEU LEU LEU	315 316 316 316 316 316	110. 877 49. 922 20. 941 1. 00 24. 80 B 108. 799 50. 772 20. 872 1. 00 21. 02 B 108. 159 49. 502 21. 184 1. 00 20. 90 B 107. 653 49. 544 22. 628 1. 00 19. 84 B 106. 866 48. 358 23. 194 1. 00 19. 46 B 107. 786 47. 157 23. 408 1. 00 18. 22 B	3 O 3 N 5 C 5 C
ATOM ATOM ATOM ATOM ATOM ATOM	8299 8300 8301 8302 8303 8304	CD2 C O N CA CB	LEU LEU ARG ARG ARG	316 316 316 317 317 317	106. 223 48. 783 24. 501 1. 00 16. 50 B 106. 995 49. 228 20. 229 1. 00 20. 90 B 106. 161 50. 098 20. 000 1. 00 22. 41 B 106. 941 48. 026 19. 666 1. 00 19. 89 B 105. 851 47. 678 18. 753 1. 00 20. 30 B 106. 154 46. 362 18. 035 1. 00 20. 73 B	C C O N C
ATOM ATOM ATOM ATOM ATOM ATOM	8305 8306 8307 8308 8309 8310	CG CD NE CZ NH1	ARG ARG ARG ARG ARG	317 317 317 317 317 317	107. 248 46. 480 16. 993 1. 00 23. 49 B 107. 524 45. 149 16. 321 1. 00 24. 95 B 108. 347 45. 314 15. 128 1. 00 25. 57 B 108. 925 44. 313 14. 476 1. 00 26. 73 B 108. 775 43. 061 14. 897 1. 00 23. 81 B	C C N C N
ATOM ATOM ATOM ATOM ATOM ATOM	8311 8312 8313 8314 8315 8316	C O N CA CB	ARG ARG ARG ARG ARG	317 317 318 318 318	104. 537 47. 545 19. 512 1. 00 19. 31 B 104. 541 47. 266 20. 713 1. 00 17. 59 B 103. 415 47. 747 18. 820 1. 00 18. 54 B 102. 117 47. 621 19. 476 1. 00 17. 04 B 100. 970 47. 781 18. 483 1. 00 17. 09 B	C O N C C
ATOM ATOM ATOM ATOM ATOM	8317 8318 8319 8320 8321	CD NE CZ NH1 NH2	ARG	318 318 318 318 318 318	99. 608 47. 794 19. 164 1. 00 17. 74 B 98. 613 48. 660 18. 414 1. 00 16. 48 B 97. 326 48. 672 19. 092 1. 00 16. 05 B 96. 320 49. 478 18. 771 1. 00 17. 02 B 96. 464 50. 342 17. 771 1. 00 13. 59 B 95. 180 49. 428 19. 460 1. 00 12. 42 B	N C N N
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8322 8323 8324 8325 8326 8327 8328	C O N CA CB CG2 CG1		318 318 319 319 319 319 319	102. 085 46. 251 20. 132 1. 00 15. 28 B 101. 569 46. 103 21. 234 1. 00 15. 74 B 102. 627 45. 251 19. 440 1. 00 15. 27 B 102. 757 43. 912 20. 007 1. 00 15. 37 B 103. 006 42. 848 18. 949 1. 00 15. 60 B 103. 268 41. 519 19. 621 1. 00 17. 64 B 101. 793 42. 732 18. 036 1. 00 15. 37 B	C O N C C
ATOM	8329	CD1		319	101. 793 42. 732 18. 036 1. 00 15. 37 B 100. 524 42. 425 18. 781 1. 00 15. 54 B	C

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					FI	G. 4	- 171	L		(Cont	inued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8330 8331 8332 8333 8334 8335 8336 8337 8341 8342 8343 8344 8345 8351 8352 8353 8354 8355 8356 8357 8358 8361 8362 8363 8364 8365 8366 8367 8368 8371 8372 8373 8374 8377 8378	O N CABCG CD NC C O N CABCG CD NC C O N CABCG CD NC C O N CABCG CD	GLN GLN GLN GLN GLN GLN ASN ASN ASN ASN ASN TYR TYR TYR TYR	319 320 320 320 320 320 320 321 321 321 321 321 322 322 322 322 322	104. 036 105. 145 103. 850 104. 923 104. 293 103. 383 102. 833 103. 544 101. 566 105. 964 106. 399 106. 382 107. 420 106. 593 105. 745 108. 658 109. 533 108. 735 109. 605 110. 766 111. 086 112. 118 111. 520 112. 557 112. 847 113. 855 110. 115 109. 240 111. 299 111. 657 111. 623 112. 602 113. 057 113. 851 113. 360 114. 672 114. 612 115. 201 115. 201 116. 487 117. 104 118. 053 118. 682 119. 851	44. 086 44. 086 44. 367 44. 693 45. 341 46. 495 47. 533 43. 663 42. 800 40. 839 40. 839 40. 839 41. 228 43. 275 43. 644 44. 733 42. 252 43. 611 43. 726 45. 149 45. 149 46. 740 47. 418 48. 638 49. 550 49. 550 50. 011 51. 243 50. 946 50. 011 51. 243 52. 280 52. 280 53. 644 545. 537 556. 946 567. 647. 648 668. 648 668. 649 669. 649. 649. 649. 649. 649. 649. 649.	20. 802 20. 257 22. 092 23. 016 24. 248 23. 863 25. 048 26. 016 24. 966 23. 437 24. 594 22. 520 22. 875	1.00 16.78 1.00 16.37 1.00 17.82 1.00 18.01 1.00 16.48 1.00 17.06 1.00 18.02 1.00 16.46 1.00 18.97 1.00 20.18 1.00 19.64 1.00 21.44 1.00 23.79 1.00 27.68	B B B B B B B B B B B B B B B B B B B	CONCCCONCONCCCONCONCCCCCCCCCCONCCCONCCCCCONCCCONCCCONCCCONCCCONCCCCONCCCCONCCCCONCCCCONCCCCONCCCCONCCCCONCCCCONCCCCONCCCCONCCCCONCCCCONCCCCCONCCCCONCCCCCONCCCCCONCCCCCONCCCCCONCCCCCONCCCCCC	
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						(Continued)
,			•	-	F I G. 4 - 172	
ATOM	8379	CE	MET	325	118.765 51.442 25.211 1.00 21.39 B	Č
ATOM	8380	C	MET	325	117. 895 51. 875 19. 782 1. 00 17. 82 B	C
ATOM	8381	0	MET	325	118. 658 51. 198 19. 082 1. 00 15. 28 B	0
ATOM	8382	N	ASP	326	117. 698 53. 175 19. 607 1. 00 18. 85 B	N
ATOM	8383	CA	ASP	326	118. 409 53. 922 18. 591 1. 00 21. 89 B	C
ATOM	8384	CB	ASP	326 326	117. 436 54. 685 17. 695 1. 00 22. 04 B	C
ATOM ATOM	8385 8386	CG	ASP ASP	326	117. 533 54. 272 16. 244 1. 00 23. 15 B 116. 800 54. 855 15. 418 1. 00 25. 35 B	C 0
ATOM	8387		ASP	326	118. 334 53. 366 15. 922 1. 00 23. 67 B	0
ATOM	8388	C	ASP	326	119. 299 54. 904 19. 327 1. 00 24. 54 B	Č
ATOM	8389	ŏ	ASP	326	118. 896 55. 494 20. 335 1. 00 25. 63 B	ŏ
ATOM	8390	Ň	ILE	327	120. 521 55. 062 18. 842 1. 00 25. 49 B	N
ATOM	8391	CA	ILE	327	121. 451 55. 986 19. 459 1. 00 27. 44 B	Ċ
ATOM	8392	CB	ILE	327	122. 713 55. 263 19. 936 1. 00 27. 10 B	C
ATOM	8393	CG2	ILE	327	123. 697 56. 264 20. 515 1. 00 27. 85 B	C
ATOM	8394	CG1	ILE	327	122. 321 54. 221 20. 984 1. 00 25. 49 B	С
ATOM	8395	CD1	ILE	327	123.476 53.506 21.594 1.00 27.60 B	C .
ATOM	8396	C	ILE	327	121. 784 57. 005 18. 395 1. 00 29. 15 B	C
ATOM	8397.	0	ILE	327	122. 357 56. 673 17. 357 1. 00 31. 19 B	0
ATOM	8398	N	CYS	328	121. 414 58. 250 18. 653 1. 00 30. 14 B	Ŋ
ATOM	8399	CA	CYS	328	121. 624 59. 298 17. 684 1. 00 31. 56 B	<u>C</u> .
ATOM	8400	C	CYS	328	122. 624 60. 356 18. 084 1. 00 32. 64 B	C
ATOM	8401	(D	CYS	328	122. 525 60. 972 19. 153 1. 00 33. 03 B	0
ATOM ATOM	8402 8403	CB SG	CYS CYS	328 328	120. 286 59. 938 17. 366 1. 00 32. 73 B	C
ATOM	8404	N	ASP	329	118. 979 58. 689 17. 154 1. 00 36. 31 B 123. 596 60. 555 17. 200 1. 00 32. 72 B	S
ATOM	8405	CA	ASP	329	123.596 60.555 17.200 1.00 32.72 B 124.639 61.542 17.406 1.00 32.74 B	N C
ATOM	8406	CB	ASP	329	125. 997 60. 975 16. 981 1. 00 34. 70 B	Č ,
ATOM	8407	CG	ASP	329	126. 480 59. 858 17. 894 1. 00 36. 73 B	č
ATOM	8408		ASP	329	127. 643 59. 431 17. 735 1. 00 38. 23 B	ŏ
ATOM	8409		ASP	329	125. 706 59. 405 18. 767 1. 00 36. 00 B	Ŏ
ATOM	8410	C	ASP	329	124. 320 62. 781 16. 588 1. 00 31. 70 B	Č
ATOM	8411	0	ASP	329	123. 767 62. 692 15. 494 1. 00 30. 70 B	0
ATOM	8412	N	TYR	330	124.662 63.940 17.129 1.00 31.69 B	N
ATOM	8413		TYR	330	124. 420 65. 191 16. 428 1. 00 33. 40 B	C
ATOM	8414	CB	TYR	330	124. 376 66. 354 17. 411 1. 00 30. 81 B	С
ATOM	8415	CG	TYR	330	124. 322 67. 693 16. 728 1. 00 29. 75 B	C
ATOM	8416	CD1	TYR	330	123.185 68.089 16.030 1.00 30.07 B	Ċ
ATOM	8417		TYR	330	123. 121 69. 326 15. 399 1. 00 30. 94 B	C
ATOM	8418		TYR	330	125. 407 68. 568 16. 777 1. 00 30. 62 B	C
ATOM	8419		TYR	330	125.356 69.814 16.150 1.00 30.16 B	C
ATOM ATOM	8420 8421	CZ OH	TYR TYR	330 330	124. 206 70. 186 15. 465 1. 00 31. 10 B 124. 122 71. 422 14. 867 1. 00 29. 92 B	C
ATOM	8422	C	TYR	330	124. 122 71. 422 14. 867 1. 00 29. 92 B 125. 523 65. 462 15. 412 1. 00 35. 09 B	0
ATOM	8423	Ö	TYR	330	126.692 65.552 15.772 1.00 36.29 B	C 0
ATOM	8424	N	ASP	331	125. 149 65. 600 14. 146 1. 00 37. 07 B	N N
ATOM	8425	CA	ASP	331	126. 123 65. 886 13. 106 1. 00 39. 50 B	C
ATOM	8426	CB	ASP	331	125. 611 65. 391 11. 756 1. 00 39. 77 B	č
ATOM	8427	ČĞ	ASP	331	126. 665 65. 464 10. 677 1. 00 40. 31 B	č
						-

	FIG. 4-173	(Continued)
ATOM 8428 OD1 ASP 33 ATOM 8429 OD2 ASP 33 ATOM 8430 C ASP 33 ATOM 8431 O ASP 33 ATOM 8432 N GLU 33 ATOM 8434 CB GLU 33 ATOM 8435 CG GLU 33 ATOM 8435 CG GLU 33 ATOM 8436 CD GLU 33 ATOM 8437 OE1 GLU 33 ATOM 8438 OE2 GLU 33 ATOM 8439 C GLU 33 ATOM 8441 N SER 333 ATOM 8441 N SER 333 ATOM 8442 CA SER 333 ATOM 8444 OG SER 333 ATOM 8445 C SER 333 ATOM 8446 O SER 333 ATOM 8446 O SER 333 ATOM 8447 N SER 334 ATOM 8448 CA SER 334 ATOM 8449 CB SER 334 ATOM 8449 CB SER 334 ATOM 8449 CB SER 334 ATOM 8450 OG SER 334 ATOM 8450 OG SER 334 ATOM 8451 C SER 334 ATOM 8450 OG SER 334 ATOM 8451 C SER 336 ATOM 8450 OG SER 334 ATOM 8450 OG SER 333 ATOM 8450 OG SER 334 ATOM 8450 OG SER 334 ATOM 8450 OG SER 333 ATOM 845	1 126. 387 65. 018 9. 543 1. 00 41. 37 1 127. 770 65. 966 10. 967 1. 00 40. 07 1 126. 355 67. 395 13. 062 1. 00 41. 15 1 125. 641 68. 126 12. 380 1. 00 40. 39 127. 358 67. 852 13. 802 1. 00 44. 16 127. 690 69. 271 13. 879 1. 00 47. 17 129. 001 69. 457 14. 646 1. 00 48. 80 129. 367 70. 901 14. 922 1. 00 51. 70 130. 451 71. 028 15. 979 1. 00 54. 56 130. 203 70. 623 17. 136 1. 00 55. 51 131. 552 71. 528 15. 658 1. 00 56. 11 127. 791 69. 941 12. 517 1. 00 47. 83 127. 518 71. 130 12. 383 1. 00 48. 69 128. 312 69. 715 10. 161 1. 00 49. 93 129. 246 68. 835 9. 327 1. 00 50. 95 130. 521 68. 723 9. 943 1. 00 50. 95 126. 514 70. 893 9. 108 1. 00 50.	B

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					FIG.	4 - 174			(Continued)
ATOM	8477		2 TRP			825 18.875	1.00 18.08	В	С
ATOM	8478		3 TRP		121.932 64.	567 19.196	1.00 15.24	В	C
ATOM	8479	CH2	2 TRP		121.798 65.	900 19.622	1.00 16.71	В	C
ATOM	8480	C	TRP		120.940 62.	487 13.188	1.00 31.57	В	C
ATOM	8481	0	TRP			167 12.482	1.00 33.23	В	0
ATOM	8482	N	ASN			712 13.347	1.00 32.12	В	N
ATOM	8483	CA	ASN		122.079 60.	426 12.691	1.00 33.02	В	C
ATOM	8484	CB	ASN				1.00 34.88	В	C
ATOM	8485	CG	ASN		122. 957 61.		1.00 38.68	В	C
ATOM	8486		ASN			306 10.251	1.00 39.82	В	0
ATOM	8487		2 ASN			845 9.669	1.00 38.06	В	N
ATOM	8488	C	ASN			294 13.693	1.00 33.48	В	C .
ATOM	8489	0	ASN			364 14.631	1.00 33.12	В	0
ATOM	8490	N	CYS		121.419 58.		1.00 33.60	В	N
ATOM	8491	CA	CYS			104 14.385	1.00 34.06	В	C
ATOM	8492	C	CYS		121. 924 55.		1.00 33.56	В	C
ATOM	8493	0	CYS		121. 135 55.		1.00 34.05	В	0
ATOM	8494	CB	CYS		120. 071 56.		1.00 34.96	В	C
ATOM	8495	SG	CYS		118. 997 58.		1.00 37.83	В	S
ATOM	8496	N	LEU		123. 211 55. (1.00 32.80	В	N
ATOM	8497	CA	LEU		123. 798 54.		1.00 33.83	В	C
ATOM	8498	CB	LEU	340	125. 303 54.		1.00 34.61	В	C
ATOM	8499	CG	LEU	340	126. 163 55. 5		1.00 34.61	В	C
ATOM	8500		LEU	340	127. 500 55. (1.00 31.70	В	С .
ATOM	8501		LEU	340	126.352 55.2		1.00 33.80	В	C
ATOM ATOM	8502	C	LEU	340	123. 152 53. 1		1.00 34.95	В	C
ATOM	8503 8504	0 N	LEU VAL	340	123.061 52.7		1.00 34.65	В	0
ATOM	8505	CA	VAL	341 341	122. 706 52. 4		1.00 35.87	В	N
ATOM	8506	CB	VAL	341	122. 093 51. 1 121. 981 50. 4		1.00 36.37	В	C
ATOM	8507		VAL	341	121. 012 49. 2		1.00 36.86	В	C
ATOM	8508		VAL	341	121.532 51.3		1.00 37.20	В	C
ATOM	8509	C	VAL	341	122. 957 50. 3		1.00 38.15 1.00 36.74	В	C
ATOM	8510	ŏ	VAL	341	122.511 49.8		1.00 30.74	B B	C 0
ATOM	8511			342	124. 200 50. 0		1.00 35.77	_	
ATOM	8512	CA	ALA	342	125. 134 49. 2		1.00 33.34	B B	N C
ATOM	8513	CB	ALA	342	126.546 49.4		1.00 34.41	В	C
ATOM	8514	Č	ALA	342	125.095 49.6		1.00 34.41	В	C
ATOM	8515	Ŏ	ALA	342	125.698 48.8		1.00 36.76	В	0
ATOM	8516	N	ARG	343	124.411 50.6		1.00 30.10	В	N
ATOM	8517	CA	ARG	343	124.303 51.0		1.00 30.81	В	C
ATOM	8518	CB	ARG	343	124.611 52.5		1.00 30.61	В	C
ATOM	8519	CG	ARG	343	126.063 52.9		1.00 34.14	В	Č
ATOM	8520	CD	ARG	343	126. 345 54. 3		1.00 33.56	В	Č
ATOM	8521	NE	ARG	343	127. 775 54. 6		1.00 33.70	В	N
ATOM	8522	CZ	ARG	343	128. 301 55. 8		1.00 34.14	В	Č
ATOM	8523	NH1		343	127. 516 56. 9		1.00 33.88	В	Ň
ATOM	8524	NH2		343	129.615 56.0		1. 00 33. 78	B	N
ATOM	8525	C	ARG	343	122.919 50.7		1.00 29.28	B	Ĉ

			•				_ :			(Coı	ntinued)
				.,	FIG	. 4 -	175				
ATOM	8526	0	ARG	343	122.586	51.143		1.00 28.30	В	0	
ATOM	8527	N	GLN	344	122. 121	50.026		1.00 28.05	В	N	
ATOM	8528	CA	GLN	344	120. 786	49.625	17. 183	1.00 28.26	·B	C	
ATOM ATOM	8529 8530	CB CG	GLN GLN	344 344		49. 238		1.00 26.68	В	C	
ATOM	8531	CD	GLN	344 344		50. 296 49. 802	15. 516 14. 399	1.00 30.39 1.00 31.50	В	C	
ATOM	8532		I GLN	344		48. 685	14. 355	1.00 31.50	B B	C 0	
ATOM	8533		2 GLN	344		50.632	13. 378	1.00 31.32	В	N	
ATOM	8534	C	GLN	344		48. 431	18. 121	1.00 28.55	В	Č	
ATOM	8535	0	GLN	344		47.515	17. 919	1.00 28.32	В	ŏ	
ATOM	8536	N	HIS	345		48.436	19.145	1.00 28.34	B	Ň	
ATOM	8537	CA	HIS	345		47. 329	20.085	1.00 28.01	В	С	
ATOM	8538	CB	HIS	345		47. 753	21.452	1.00 28.88	В	C	
ATOM ATOM	8539 8540	CC	HIS HIS	345 345		48.079	21.443	1.00 27.88	B	C	
ATOM	8541		HIS	$\frac{345}{345}$		47. 279 49. 361	21. 516 21. 270	1.00 26.67 1.00 28.37	В	C	
ATOM	8542		HIS	345		49. 337	21. 270	1.00 28.37	B B	N C	
ATOM	8543		HIS	345		48. 086	21. 381	1.00 28.63	В	N	•
ATOM	8544	C	HIS	345		46. 799	20. 215	1.00 27.76	B	Ċ	
ATOM	8545	0	HIS	345	117.659	47. 508	20.625	1.00 30.01	B	Ŏ	
ATOM	8546	N	ILE	346		45. 538	19.849	1.00 26.83	В	N	
ATOM ATOM	8547 8548	CA CB	ILE ILE	346		44. 897	19.899	1.00 25.72	В	C	
ATOM	8549		ILE	$\begin{array}{c} 346 \\ 346 \end{array}$		43.842	18. 791	1.00 25.56	В	C	
ATOM	8550		ILE	346		43. 114 44. 517	18. 919 17. 422	1.00 26.17 1.00 26.62	В	C	
ATOM	8551		ILE	346		43. 544	16. 263	1.00 26.02	B B	C	
ATOM	8552	C ·	ILE	346		44. 218	21. 228	1.00 26.11	В	C	
ATOM	8553	0	ILE	346		43. 558	21.776	1.00 25.75	B	ŏ	
ATOM	8554	N	GLU	347		44.396	21.746	1.00 26.23	В	N	
ATOM ATOM	8555	CA	GLU	347		43. 767	22. 994	1.00 25.82	В	C	
ATOM	8556 8557	CB CG	GLU GLU	347 347		14.777	24. 134	1.00 25.51	В	C	
ATOM	8558	CD	GLU	347		14. 118 15. 094	25. 505	1.00 28.20	В	C	
ATOM	8559		GLU	347		16. 208	26. 652 26. 592	1.00 29.16 1.00 29.18	B B	C .	
ATOM	8560		GLU	347		14.736	27. 628	1.00 23.16	В	0	
ATOM	8561	C	GLU	347		13. 172	22. 799	1.00 26.44	В	Č	
ATOM	8562	0	GLU	347	112.919 4	13.889	22.495	1.00 26.00	$\tilde{\mathbf{B}}$	Ŏ	
ATOM	8563	N	MET	348		11.858	22.957	1.00 26.58	В	N	
ATOM ATOM	8564 8565	CA CB	MET	348		1.181	22. 807	1.00 27.90	В	. C	
ATOM	8566	CG	MET MET	348 348		0.767	21.345	1.00 30.41	В	C	
ATOM	8567	SD	MET	348		0.132 8.420	20.660 21.117	1.00 34.65	. B	C	
ATOM	8568	CE	MET	348		7. 597	19. 804	1.00 42.21 1.00 38.96	B B	S C	
ATOM	8569	C	MET	348		9. 980	23. 732	1.00 26.60	В	C	
ATOM	8570	0	MET	348		9. 472	24. 247	1.00 26.08	В	ŏ	
ATOM	8571	N	SER	349	111.135 3	9.549	23.950	1.00 23.99	B	Ň	
ATOM	8572	CA	SER	349		8. 423	24. 812	1.00 21.78	В	C	
ATOM ATOM	8573 8574	CB OG	SER SER	349 349		8.894		1.00 20.79	В	C	
* 11 Oli1	0017	UU	DD1/			7. 809		1.00 21.42	·B	0	
				30	JBSTITUTE S	חבבו (ו	KULE 26)				

f-ca					FIG. 4-176	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8610	C O N CA CB OG1 CG2 C O N CA CB CG CCD2 CE2 CC3 CCD1 NE1 CZ2 CC3 CCH2 C O N CA	THRIRING THRICK	349 349 350 350 350 351 351 351 351 352 352 353 353 353 353 353 353 353 353	FIG. 4 - 176 110.084 37.387 24.005 1.00 21.88 B 109.274 37.739 23.154 1.00 23.74 B 110.351 36.112 24.264 1.00 21.76 B 109.654 35.033 23.571 1.00 23.08 B 110.603 33.882 23.214 1.00 22.77 B 111.310 33.483 24.391 1.00 25.37 B 111.583 34.299 22.152 1.00 22.93 B 108.561 34.453 24.475 1.00 22.93 B 107.732 33.650 24.035 1.00 20.70 B 108.564 34.871 25.737 1.00 22.30 B 107.601 34.366 26.703 1.00 22.35 B 108.332 33.796 27.932 1.00 23.36 B 108.989 34.859 28.635 1.00 25.67 B 109.378 32.781 27.493 1.00 25.67 B 106.839 36.668 26.918 1.00 21.07 B 105.562 35.031 27.760 1.00 21.07 B 105.562 35.031 27.760 1.00 20.87 B 106.839 36.668 26.918 1.00 19.83 B 105.894 37.692 27.325 1.00 19.36 B 106.182 39.027 26.672 1.00 18.63 B 105.913 40.109 27.397 1.00 15.30 B 105.195 42.451 27.587 1.00 13.08 B 105.564 34.387 29.877 1.00 7.79 B 104.739 41.684 31.233 1.00 8.17 B 104.479 41.387 29.877 1.00 7.79 B 104.739 41.684 31.233 1.00 8.17 B 104.739 41.684 31.233 1.00 8.17 B 105.546 42.791 31.265 1.00 10.10 B 104.217 40.921 32.281 1.00 10.66 B 103.149 39.524 30.625 1.00 10.10 B 104.217 40.921 32.281 1.00 10.66 B 103.149 39.524 30.625 1.00 10.10 B 104.217 40.921 32.281 1.00 10.66 B 103.149 39.524 30.625 1.00 10.40 B 103.426 39.848 31.958 1.00 9.81 B 107.594 41.796 27.264 1.00 15.80 B 108.247 40.999 27.931 1.00 13.84 B 109.464 43.338 27.140 1.00 13.65 B	Continued) C O N C C O C C O N C C C O N C C C C
ATOM ATOM ATOM	8611 8612 8613	CB	VAL VAL	354 354 354	110. 135	C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8614 8615 8616 8617 8618 8619 8620 8621 8622	COLUMN CA CA CB CG	VAL VAL GLY GLY GLY GLY ARG ARG	354 354 355 355 355 356 356 356 356	110. 264 43. 163 24. 751 1. 00 12. 49 B 109. 486 44. 248 28. 368 1. 00 13. 83 B 108. 716 45. 197 28. 456 1. 00 13. 93 B 110. 373 43. 957 29. 313 1. 00 14. 87 B 110. 467 44. 769 30. 519 1. 00 16. 09 B 109. 333 44. 554 31. 513 1. 00 16. 34 B 108. 347 43. 877 31. 206 1. 00 18. 25 B 109. 456 45. 126 32. 706 1. 00 15. 16 B 108. 404 44. 953 33. 701 1. 00 16. 32 B 108. 856 45. 494 35. 066 1. 00 14. 18 B 110. 001 44. 668 35. 667 1. 00 13. 44 B	C C O N C C O N C C

					FIG. 4-177	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8624 8625 8626 8627 8628 8631 8632 8633 8633 8633 8634 8635 8636 8644 8645 8645 8655 8655 8655 8655 8665 866	NE CZ NH NH: C O N CA CB CCD CC2 CC O N CA CB CCD NE CC NH CCZ NH I	ARG ARG HEELEHEELEHEELE ARG ARG ARG PROOPRO PROPRO PROPROPRO PROPRO PROPROPRO PROPRO P	356 356 356 356 356 356 357 357 357 357 357 357 357 357 357 357	FIG. 4 - 177 110.169 44.878 37.151 1.00 14.42 B 111.546 45.211 37.511 1.00 18.65 B 112.457 44.341 37.935 1.00 20.17 B 112.156 43.055 38.065 1.00 22.71 B 113.674 44.765 38.242 1.00 18.93 B 107.111 45.607 33.209 1.00 16.01 B 106.100 44.924 33.066 1.00 16.29 B 107.140 46.911 32.945 1.00 15.89 B 105.967 47.603 32.402 1.00 16.40 B 105.418 48.660 33.366 1.00 11.21 B 104.753 48.083 34.573 1.00 8.48 B 105.467 47.878 35.748 1.00 5.58 B 103.407 47.711 34.531 1.00 8.57 B 104.846 47.309 36.867 1.00 5.98 B 102.777 47.136 35.648 1.00 4.59 B 105.476 48.638 30.287 1.00 18.69 B 105.476 48.638 30.287 1.00 19.12 B 107.648 48.377 30.840 1.00 19.12 B 108.188 48.953 29.612 1.00 19.47 B 107.826 50.439 29.499 1.00 19.02 B 108.451 51.346 30.559 1.00 19.99 B 108.074 52.820 30.338 1.00 22.48 B 109.204 54.890 31.117 1.00 24.69 B 109.204 54.890 31.117 1.00 24.69 B 109.304 55.358 29.875 1.00 21.57 B 109.304 55.358 29.875 1.00 21.14 B 109.696 55.603 32.121 1.00 24.69 B 109.707 48.784 29.646 1.00 20.57 B 110.302 48.704 30.722 1.00 24.13 B 111.816 48.564 28.411 1.00 20.48 B 112.137 48.916 26.959 1.00 19.85 B 112.221 50.683 29.465 1.00 20.23 B 113.474 48.953 30.163 1.00 19.33 B 114.212 49.725 31.160 1.00 18.75 B 115.122 48.806 31.968 1.00 20.74 B 115.410 50.806 29.382 1.00 17.99 B 115.394 51.824 31.393 1.00 18.96 B	Continued) C N C N C N C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM	8667 8668 8669 8670 8671 8672	CA CB CG	GLU GLU GLU GLU GLU	361 361 361 361 361 361	116. 199 52. 970 30. 978 1. 00 18. 11 B 115. 982 54. 159 31. 919 1. 00 16. 34 B 116. 654 54. 007 33. 269 1. 00 21. 67 B 115. 743 53. 431 34. 342 1. 00 27. 42 B 115. 067 52. 408 34. 091 1. 00 28. 62 B	C C C C
					115.710 54.009 35.453 1.00 31.11 B	0 .

					FIG	S. 4-	178			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	8673 8674 8675 8676 8677 8678 8679 8680 8681 8682 8683 8684 8685 8686 8687 8688 8690 8691 8692 8693 8694 8695 8696 8697 8698 8701 8702 8703 8704 8705 8706 8707 8708 8709 8710 8711 8712	ND1 CE1 NE2 C O N CA CB CCD1 CE2 CZ C O N CA CB OCA CB CCD1 CE2 CZ C O N CA CB CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD1 CCD2 CCD2	GLUOOROOROORISSSSSSSSSSSSSSSSSSSSSSSSSSSSS	361 362 362 362 363 363 363 363 363 363 364 364 364 364	117. 674 118. 118 118. 449 118. 027 119. 879 120. 207 119. 362 120. 601 120. 096 121. 768 122. 550 122. 626 121. 324 120. 156 121. 111 119. 869 119. 267 123. 942 124. 833 124. 110 125. 371 125. 188 124. 368 122. 975 124. 989 122. 216 124. 225 122. 837 126. 531 126. 341 127. 735 128. 967 130. 132 130. 257 129. 890 129. 312 128. 662 130. 329 130. 740	52. 595 51. 870 53. 079 53. 805 52. 772 52. 916 54. 086 54. 911 53. 448 54. 374 53. 875 53. 965 53. 801 54. 731 54. 499 53. 691 55. 520 56. 319 56. 339 55. 770 55. 816 55. 242 56. 127 56. 638 57. 638 58. 163 58. 163 59. 544	31. 007 31. 888 30. 030 28. 817 29. 985 28. 505 28. 121 30. 832 31. 021 31. 353 32. 164 33. 603 34. 333 34. 158 35. 799 35. 081 31. 551 31. 806 30. 723 30. 043 28. 944 27. 777 27. 826 26. 656 26. 781 25. 607 25. 679 30. 942 32. 050 30. 448 31. 159 30. 697 29. 275 31. 069 30. 697 29. 275 31. 304	1. 00 16. 97 1. 00 16. 23 1. 00 16. 09 1. 00 13. 66 1. 00 15. 32 1. 00 12. 78 1. 00 12. 78 1. 00 17. 05 1. 00 17. 21 1. 00 18. 58 1. 00 19. 36 1. 00 19. 36 1. 00 19. 36 1. 00 19. 36 1. 00 19. 50 1. 00 19. 50 1. 00 19. 73 1. 00 19. 73 1. 00 19. 25 1. 00 17. 71 1. 00 15. 99 1. 00 12. 83 1. 00 19. 25 1. 00 17. 71 1. 00 15. 99 1. 00 12. 83 1. 00 12. 86 1. 00 8. 09 1. 00 12. 83 1. 00 12. 86 1. 00 8. 09 1. 00 17. 71 1. 00 18. 72 1. 00 17. 73 1. 00 17. 73 1. 00 17. 73 1. 00 17. 73 1. 00 17. 73 1. 00 17. 73 1. 00 17. 73 1. 00 17. 73 1. 00 22. 16 1. 00 13. 36 1. 00 20. 48 1. 00 20. 68 1. 00 22. 60 1. 00 25. 75 1. 00 29. 32	B B B B B B B B B B B B B B B B B B B	CONCCCCONCCCNCNCONCCCCCCCCONCCOOCCONC
ATOM ATOM ATOM ATOM	8712 8713 8714 8715	CB CG CD1 CD2	LEU LEU LEU	366 366 366 366	132. 053 132. 172 132. 442 133. 316	59. 831 59. 429 57. 920 60. 210	32. 039 33. 516 33. 631 34. 162	1.00 29.32 1.00 34.01 1.00 33.57 1.00 34.78	В В В	C C C
ATOM ATOM ATOM ATOM ATOM ATOM	8715 8716 8717 8718 8719 8720	CD2 C O N CA CB	LEU LEU LEU ASP ASP ASP	366 366 367 367	130. 909 130. 317 131. 709 131. 964	59. 900 60. 871 59. 115 59. 369	29. 824 29. 349 29. 102 27. 682	1.00 26.20 1.00 26.53 1.00 24.26 1.00 23.63	B B B	C O N C
ATOM	8721	CG	ASP	367 367		58. 636 57. 158	27. 214 27. 582	1.00 23.47 1.00 25.27	B B	C

ATOM 8722 OD1 ASP 367 132.158 56.515 27.507 1.00 24.35 B O ATOM 8723 OD2 ASP 367 134.311 56.634 27.935 1.00 25.99 B O ATOM 8724 C ASP 367 130.810 58.990 26.767 1.00 22.76 B C ATOM 8725 O ASP 367 130.848 59.261 25.568 1.00 24.31 B O ATOM 8726 N GLY 368 129.795 58.348 27.330 1.00 20.91 B N ATOM 8727 CA GLY 368 128.646 57.950 26.547 1.00 18.80 B C ATOM 8728 C GLY 368 128.912 56.843 25.550 1.00 19.81 B C ATOM 8729 0 GLY 368 128.059 56.563 24.700 1.00 19.20 B <t< th=""><th></th></t<>	
ATOM 8731 CA ASN 369 130.073 56.198 25.643 1.00 19.20 B N ATOM 8732 CB ASN 369 131.907 54.986 24.526 1.00 19.60 B C ATOM 8733 CG ASN 369 132.519 56.217 23.921 1.00 21.94 B C ATOM 8734 OD1 ASN 369 132.005 56.757 22.945 1.00 25.32 B O ATOM 8735 ND2 ASN 369 132.005 56.757 22.945 1.00 23.16 B N ATOM 8736 C ASN 369 132.005 56.757 22.945 1.00 23.16 B N ATOM 8736 C ASN 369 129.828 53.760 25.090 1.00 18.53 B C ATOM 8737 O ASN 369 129.828 53.760 25.090 1.00 18.53 B C ATOM 8738 N SER 370 129.420 53.608 26.346 1.00 18.61 B N ATOM 8738 N SER 370 129.420 53.608 26.346 1.00 18.61 B N ATOM 8739 CA SER 370 129.934 51.447 27.430 1.00 20.45 B C ATOM 8740 CB SER 370 129.934 51.447 27.430 1.00 20.45 B C ATOM 8741 OG SER 370 127.746 52.621 27.7829 1.00 18.95 B C ATOM 8742 C SER 370 127.746 52.621 27.7829 1.00 18.95 B C ATOM 8744 N PHE 371 127.009 51.583 28.209 1.00 18.63 B N ATOM 8745 CA PHE 371 125.931 51.763 29.168 1.00 19.79 B C ATOM 8746 CB PHE 371 124.762 52.516 28.512 1.00 19.79 B C ATOM 8748 CD PHE 371 124.468 51.766 27.398 1.00 18.63 B N ATOM 8749 CD2 PHE 371 124.483 51.98 25.059 1.00 18.69 B C ATOM 8749 CD2 PHE 371 124.532 51.874 25.093 1.00 18.63 B N ATOM 8745 CA PHE 371 124.532 51.874 25.093 1.00 18.69 B C ATOM 8746 CB PHE 371 124.532 51.874 25.093 1.00 18.69 B C ATOM 8747 CG PHE 371 124.532 51.874 25.093 1.00 18.69 B C ATOM 8748 CD1 PHE 371 124.532 51.874 25.093 1.00 18.69 B C ATOM 8750 CE1 PHE 371 122.340 50.255 26.631 1.00 19.79 B C ATOM 8751 CE2 PHE 371 122.340 50.255 26.631 1.00 18.78 B C ATOM 8752 CZ PHE 371 122.340 50.255 26.631 1.00 18.78 B C ATOM 8753 C PHE 371 122.448 49.50 33.290 1.00 18.89 B C ATOM 8756 C PHE 371 122.449 49.491 31.703 1.00 18.79 B C ATOM 8757 CB TYR 372 124.844 49.491 31.703 1.00 18.89 B C ATOM 8760 CE1 TYR 372 124.897 49.557 33.3491 1.00 18.89 B C ATOM 8760 CE1 TYR 372 124.897 49.558 31.620 1.00 18.89 B C ATOM 8760 CE1 TYR 372 124.899 49.500 33.290 1.00 18.89 B C ATOM 8766 O TYR 372 124.899 49.500 33.491 1.00 18.89 B C ATOM 8766 O TYR 372 128.266 48.257 33.498 1.00 18.88 B C ATOM 8766 O TYR	
ATOM 8768 CA LYS 373 120.647 48.340 31.299 1.00 18.51 B C ATOM 8769 CB LYS 373 120.285 48.376 29.809 1.00 17.90 B C ATOM 8770 CG LYS 373 118.809 48.581 29.485 1.00 21.01 B C	

						. .				(Continued)
					F I (3. 4 -	180			
ATOM	8771	CD	LYS	373	118. 593	48.627	27. 969	1.00 21.40	В	С
ATOM	8772	CE	LYS	373	117. 248	49. 238	27. 563	1.00 21.40	В	Č
ATOM	8773	NZ	LYS	373	116.053	48. 389	27. 855	1.00 21.07	В	N
		C	LYS	373	120. 128	47. 049	31. 928	1.00 21.38	В	C
ATOM ATOM	8774 8775	0	LYS	373	120. 126	45. 980	31. 712	1.00 18.48	В	0
ATOM	8776	N	ILE	374	119.056	47. 150	32. 709	1.00 10.40	В	N
ATOM	8777	CA	ILE	374	118. 474	45. 972	33. 332	1.00 15.88	В	
ATOM	8778	CB	ILE	374	117. 557	46. 339	34. 526	1.00 14.58	, В	C C
ATOM	8779		ILE	374	116. 955	45. 076	35. 130	1.00 12.18	В	Č
ATOM	8780		ILE	374	118.348	47. 101	35. 591	1.00 15.07	В	. Č
ATOM	8781		ILE	374	117. 517	47. 505	36. 809	1.00 13.03	B	Č
ATOM	8782	CDI	ILE	374	117.618	45. 244	32. 303	1.00 16.94	В	Č
ATOM	8783	Õ	ILE	374	116.649	45. 803	31. 795	1.00 17.41	В	ŏ
ATOM	8784	N	ILE	375	117. 977	44.008	31. 978	1.00 18.50	В	Ň
ATOM	8785	CA	ILE	375	117. 178	43. 226	31.033	1.00 19.71	B	Č
ATOM	8786	CB	ILE	375	117. 842	43.117	29.625	1.00 19.62	B	č
ATOM	8787		ILE	375	118. 128	44. 496	29.070	1.00 19.13	В	C C C
ATOM	8788		ILE	375	119. 128	42. 298	29. 706	1.00 21.23	B	Č
ATOM	8789		ILE	375	119.824	42.129	28. 373	1.00 23.06	$\tilde{\mathbf{B}}$	Č
ATOM	8790	Č	ILE	375	116. 984	41.815	31.579	1.00 20.44	B	Č
ATOM	8791	0	ILE	375	117.735	41.356	32.443	1.00 20.03	В	0
ATOM	8792	N	SER	376	115.968	41.128	31.078	1.00 21.14	В	N
ATOM	8793	CA	SER	376	115.705	39.771	31.516	1.00 21.95	В	C
ATOM	8794	CB	SER	376	114. 347	39.318	31.003	1.00 21.55	В	C
ATOM .	8795	0G	SER	376	114.026	38.054	31.539	1.00 25.40	В	0
ATOM	8796	C	SER	376	116.808	38.899	30.936	1.00 23.06	В	C
ATOM	8797	0	SER	376	117. 236	39. 127	29.807	1.00 24.16	В	0
ATOM	8798	N	ASN	377	117. 281	37.914	31.698	1.00 24.67	В	N
ATOM	8799	CA	ASN	377	118. 358	37.053	31. 218	1.00 25.07	В	C
ATOM	8800	CB	ASN	377	119. 438	36. 891	32.302	1.00 23.49	В	C
ATOM	8801	CG	ASN	377	119.010	35. 971	33. 444	1.00 23.86	В	C
ATOM	8802		ASN	377	117. 951	35.340	33. 397	1.00 23.70	В	0
ATOM	8803		ASN	377	119. 848	35. 884	34. 474	1.00 20.11	В	N
ATOM	8804			377	117. 897			1.00 26.79	В	С
ATOM	8805	0	ASN	377	116. 706	35. 382	30.699	1.00 28.58	В	0
ATOM	8806	N	GLU	378	118. 861	34.856	30. 353	1.00 29.97	В	N
ATOM	8807	CA	GLU	378	118.608	33. 504	29. 871	1.00 33.15	В	C
ATOM	8808	CB	GLU	378	119. 914	32.716	29.870	1.00 37.08	В	C
ATOM	8809	CG	GLU	378	120. 695	32.870	31. 181	1.00 43.78	В	C · .
ATOM	8810	CD	GLU	378	121.681	31.740	31.427	1.00 46.56	В	C
ATOM	8811		GLU	378	121. 225	30.613	31. 725	1.00 47.52	В	0
ATOM	8812		GLU	378	122. 906	31.981	31. 321	1.00 47.91	В	0
ATOM	8813	C	GLU	378	117.588	32.760	30. 722	1.00 33.63	В	C
ATOM	8814	0 N	GLU	378	116.685	32. 113	30. 192	1.00 35.16	В	0
ATOM	8815	N CA	GLU GLU	379 370	117.740	32. 842	32. 041	1.00 32.70	В	N
ATOM ATOM	8816 8817	CA CB	GLU	379 379	116.831	32.160	32. 953	1.00 30.44	B B	C
ATOM	8818	CG	GLU	379	117. 549	31.806 30.323	34. 256	1.00 34.46	В	C
ATOM	8819	CD	GLU	379	117. 845		34. 412	1.00 39.45 1.00 43.32	В	C C
VI OM	0013	Ųν	and	010	116. 577	29.475	34.492	1.00 40.04	ע	U

	FIG. 4-182	(Continued)
ATOM 8869 N CYS 385 ATOM 8870 CA CYS 385 ATOM 8871 C CYS 385 ATOM 8872 O CYS 385 ATOM 8873 CB CYS 385 ATOM 8874 SG CYS 385 ATOM 8875 N TYR 386 ATOM 8876 CA TYR 386 ATOM 8877 CB TYR 386 ATOM 8877 CB TYR 386 ATOM 8878 CG TYR 386 ATOM 8887 CG TYR 386 ATOM 8881 CD2 TYR 386 ATOM 8881 CD2 TYR 386 ATOM 8882 CE2 TYR 386 ATOM 8883 CZ TYR 386 ATOM 8884 OH TYR 386 ATOM 8885 C TYR 386 ATOM 8885 C TYR 386 ATOM 8886 O TYR 386 ATOM 8887 N PHE 387 ATOM 8888 CA PHE 387 ATOM 8889 CB PHE 387 ATOM 8890 CG PHE 387 ATOM 8891 CD1 PHE 387 ATOM 8891 CD1 PHE 387 ATOM 8892 CD2 PHE 387 ATOM 8893 CE1 PHE 387 ATOM 8894 CE2 PHE 387 ATOM 8895 C PHE 387 ATOM 8890 CG PHE 387 ATOM 8890 CB PHE 387 ATOM 8891 CD1 PHE 387 ATOM 8890 CB CP PHE 387 ATOM 8890 CB CP PHE 387 ATOM 8890 CB CP PHE 387 ATOM 8890 CB CD RHE 387 ATOM 8890 CB GLN 388 ATOM 8901 CG GLN 388 ATOM 8901 CG GLN 388 ATOM 8902 CD GLN 388 ATOM 8903 OE1 GLN 388 ATOM 8904 NE2 GLN 388 ATOM 8907 N ILE 389 ATOM 8909 CB ILE 389 ATOM 8901 CG2 ILE 389 ATOM 8911 CG1 ILE 389 ATOM 8911 CG1 ILE 389 ATOM 8911 CG1 ILE 389 ATOM 8912 CD1 ILE 389 ATOM 8913 C ILE 389 ATOM 8914 O ILE 389 ATOM 8915 N ASP 390 ATOM 8916 CA ASP 390	122. 956 44. 485 32. 446 1. 00 19. 06 123. 812 44. 063 31. 340 1. 00 20. 78 124. 628 45. 266 30. 868 1. 00 19. 29 124. 115 46. 376 30. 775 1. 00 19. 30	B B C C C C C C C C C C C C C C C C C C
ATOM 8917 CB ASP 390	130. 478 50. 005 18. 365 1. 00 45. 79	B C

ATOM 8918 CG ASP 390 130.576 51.416 17.816 1.00 49.16 B C ATOM 8919 OD1 ASP 390 129.879 51.713 16.819 1.00 50.13 B O ATOM 8921 C ASP 390 131.349 52.227 18.372 1.00 50.30 B O ATOM 8922 O ASP 390 128.887 48.106 18.675 1.00 44.93 B C ATOM 8922 O ASP 390 128.887 48.106 18.675 1.00 44.93 B C ATOM 8922 C ASP 390 128.887 48.106 18.675 1.00 44.93 B C ATOM 8923 N LYS 391 129.981 47.427 19.798 1.00 47.19 B O ATOM 8924 CA LYS 391 129.981 47.427 19.798 1.00 45.32 B N ATOM 8925 CB LYS 391 129.981 45.409 20.818 1.00 45.81 B C ATOM 8926 CG LYS 391 131.416 45.724 20.407 1.00 51.34 B C ATOM 8927 CD LYS 391 131.416 45.724 20.407 1.00 55.03 B C ATOM 8928 CE LYS 391 133.816 45.911 21.112 1.00 55.62 B C ATOM 8930 C LYS 391 134.822 45.719 22.192 1.00 56.68 B N ATOM 8930 C LYS 391 127.550 45.535 20.163 1.00 44.97 B N ATOM 8931 O LYS 391 126.857 46.191 20.942 1.00 46.28 B O ATOM 8932 N LYS 392 125.772 43.916 19.782 1.00 44.97 B N ATOM 8933 C LYS 392 125.772 43.916 19.782 1.00 44.97 B N ATOM 8934 CB LYS 392 125.772 43.916 19.782 1.00 46.28 B O ATOM 8936 CD LYS 392 125.772 43.916 19.782 1.00 44.97 B N ATOM 8937 C LYS 392 125.772 43.916 19.782 1.00 44.97 B N ATOM 8938 C LYS 392 125.772 43.916 19.782 1.00 44.97 B N ATOM 8936 CD LYS 392 125.772 43.916 19.782 1.00 45.02 B C ATOM 8937 C LYS 392 125.772 43.916 19.782 1.00 44.97 B N ATOM 8938 C LYS 392 125.772 43.916 19.782 1.00 44.97 B N ATOM 8936 CD LYS 392 125.772 43.916 19.782 1.00 44.97 B N ATOM 8937 C LYS 392 125.772 43.916 19.782 1.00 44.90 B C ATOM 8938 NZ LYS 392 125.772 43.916 19.782 1.00 44.91 B C ATOM 8939 C LYS 392 125.772 43.916 19.782 1.00 44.91 B C ATOM 8938 C LYS 392 125.772 43.916 19.782 1.00 44.90 B C ATOM 8939 C LYS 392 125.772 43.916 19.782 1.00 49.00 B C ATOM 8938 NZ LYS 392 125.772 43.916 19.782 1.00 49.00 B C ATOM 8937 C LYS 392 125.772 43.916 19.782 1.00 49.27 B N ATOM 8948 O LYS 393 126.679 42.486 22.470 1.00 41.43 B C ATOM 8949 O LYS 393 126.679 42.486 22.662 1.00 40.21 B C ATOM 8940 O LYS 393 126.685 42.158 2.953 1.00 43.49 B O ATOM 8947 C ASP 393 127.889 40.268 22.	ed)
ATOM 8919 OD1 ASP 390 129.879 51.713 16.819 1.00 50.13 B 0 ATOM 8920 OD2 ASP 390 131.349 52.227 18.372 1.00 50.30 B 0 ATOM 8921 C ASP 390 128.887 48.106 18.675 1.00 44.93 B C ATOM 8922 O ASP 390 128.887 48.106 18.675 1.00 44.93 B C ATOM 8923 N LYS 391 129.081 47.427 19.798 1.00 45.32 B N ATOM 8924 CA LYS 391 129.081 47.427 19.798 1.00 45.32 B N ATOM 8925 CB LYS 391 129.981 45.409 20.818 1.00 45.91 B C ATOM 8926 CG LYS 391 131.416 45.724 20.407 1.00 51.34 B C ATOM 8927 CD LYS 391 132.428 45.397 21.494 1.00 55.03 B C ATOM 8928 CE LYS 391 133.816 45.917 21.112 1.00 55.62 B C ATOM 8929 NZ LYS 391 134.822 45.719 22.192 1.00 56.68 B N ATOM 8930 C LYS 391 127.550 45.535 20.163 1.00 45.76 B C ATOM 8931 O LYS 391 126.857 46.191 20.942 1.00 44.97 B N ATOM 8932 N LYS 392 127.125 44.419 19.576 1.00 44.97 B N ATOM 8934 CB LYS 392 125.772 43.916 19.782 1.00 44.97 B N ATOM 8935 CG LYS 392 125.218 43.382 18.458 1.00 44.97 B N ATOM 8936 CD LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8937 CE LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8938 NZ LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8939 C LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8939 C LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8939 C LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8939 C LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8937 CE LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8938 NZ LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8939 C LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8939 C LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8940 0 LYS 392 124.282 43.970 16.186 1.00 50.10 B C ATOM 8941 N ASP 393 126.579 42.446 21.555 1.00 41.92 B N ATOM 8944 CG ASP 393 126.685 42.512 21.134 1.00 41.43 B C ATOM 8945 OD1 ASP 393 126.685 42.158 23.953 1.00 43.49 B O ATOM 8946 OD2 ASP 393 126.685 42.158 23.953 1.00 43.49 B O ATOM 8947 C ASP 393 126.685 42.158 23.953 1.00 43.49 B O ATOM 8948 OD ASP 393 126.685 42.158 23.953 1.00 43.49 B O	
ATOM 8949 N CYS 394 125.678 42.252 24.816 1.00 35.47 B N ATOM 8950 CA CYS 394 125.882 42.870 26.117 1.00 32.02 B C ATOM 8951 C CYS 394 126.374 41.796 27.069 1.00 29.62 B C ATOM 8952 0 CYS 394 126.248 40.608 26.787 1.00 29.41 B 0 ATOM 8953 CB CYS 394 124.586 43.491 26.639 1.00 31.92 B C ATOM 8954 SG CYS 394 123.354 42.328 27.301 1.00 33.67 B S ATOM 8955 N THR 395 126.938 42.215 28.193 1.00 26.53 B N	-
ATOM 8957 CB THR 395 128.964 41.493 29.358 1.00 23.30 B C ATOM 8958 OG1 THR 395 129.627 41.265 28.115 1.00 25.56 B O	
ATOM 8959 CG2 THR 395 129.518 40.542 30.397. 1.00 22.48 B C ATOM 8960 C THR 395 126.784 41.448 30.519 1.00 22.20 B C ATOM 8961 O THR 395 126.707 42.556 31.035 1.00 23.25 B O ATOM 8962 N PHE 396 126.300 40.354 31.095 1.00 19.02 B N	
ATOM 8963 CA PHE 396 125.658 40.444 32.396 1.00 18.94 B C ATOM 8964 CB PHE 396 124.794 39.206 32.652 1.00 17.62 B C ATOM 8965 CG PHE 396 123.486 39.225 31.918 1.00 19.32 B C	
ATOM 8966 CD1 PHE 396 122.477 40.112 32.290 1.00 20.73 B C	

					FIG	. 4 -	184			(Continued)
ATOM ATOM ATOM	8967 8968 8969	CE1	PHE PHE PHE	396 396 396	123. 265 121. 267 122. 062	38. 378 40. 157 38. 411	30. 837 31. 593 30. 130	1.00 19.67 1.00 21.82 1.00 20.02	B B B	C C C
ATOM ATOM ATOM	8970 8971 8972	CZ C O	PHE PHE PHE	396 396 396	121. 057 126. 712 127. 703	39. 303 40. 596 39. 866	30. 507 33. 488 33. 516	1.00 22.36 1.00 19.09 1.00 21.70	В В В	C C C O
ATOM ATOM ATOM	8973 8974 8975	N CA CB	ILE ILE ILE	397 397 397	126. 511 127. 454 127. 819	41.559 41.774 43.240	34. 380 35. 460 35. 566	1.00 17.18 1.00 14.91 1.00 14.47	B B B	N
ATOM ATOM ATOM	8976 8977 8978	CG2 CG1	ILE ILE ILE	397 397 397	128. 181 126. 644 126. 993	43. 762 44. 036 45. 472	34. 192 36. 135 36. 449	1.00 14.09 1.00 13.14 1.00 11.32	B B B	C C C C C
ATOM ATOM ATOM	8979 8980 8981	C O N	ILE ILE THR	397 397 398	126. 885 127. 543 125. 651	41. 287 41. 376 40. 790	36. 791 37. 833 36. 753	1.00 16.82 1.00 18.48 1.00 15.47	B B B	C O N
ATOM ATOM ATOM	8982 8983 8984	CA CB	THR THR THR	398 398 398	125. 000 124. 049 122. 968	40. 241 41. 255 41. 627	37. 937 38. 652 37. 784	1.00 14.86 1.00 14.72 1.00 13.55	B B B	C C O
ATOM ATOM ATOM	8985 8986 8987	C 0	THR THR THR	398 398 398	124. 812 124. 185 123. 805	42. 476 39. 040 38. 942	39. 083 37. 490 36. 323	1.00 13.88 1.00 15.72 1.00 15.48	B B B	C C O
ATOM ATOM ATOM	8988 8989 8990	N CA CB	LYS LYS LYS	399 399 399	123. 915 123. 147 124. 026	38. 127 36. 935 35. 960	38. 416 38. 094 37. 314	1.00 17.12 1.00 18.19 1.00 20.96	B B B	N C C C C
ATOM ATOM ATOM	8991 8992 8993	CE	LYS LYS LYS	399 399 399	125. 322 125. 970 127. 055	35. 630 34. 380 33. 860	38. 023 37. 458 38. 402	1.00 24.93 1.00 29.93 1.00 32.81	В В В	C
ATOM ATOM ATOM ATOM	8994 8995 8996 8997	NZ C O N	LYS LYS LYS GLY	399 399 399 400	128. 082 122. 616 123. 041	34. 904 36. 259 36. 571	38. 703 39. 354 40. 465	1.00 34.86 1.00 17.75 1.00 18.35	B B B	N C O
ATOM ATOM ATOM ATOM	8998 8999 9000	CA C O	GLY GLY GLY	400 400 400 400	121. 684 121. 131 119. 616 118. 979	35. 331 34. 640 34. 629 35. 360	39. 181 40. 327 40. 320 39. 551	1.00 16.55 1.00 17.62 1.00 19.66	B B B	N C C
ATOM ATOM ATOM	9001 9002 9003		THR THR THR	401 401 401	119. 028 117. 582 117. 125	33. 797 33. 708 32. 323		1.00 22.36 1.00 18.45 1.00 17.93 1.00 17.98	B B B	0 N C C
ATOM ATOM ATOM	9004 9005 9006	0G1	THR THR THR	401 401 401	117. 653 117. 607 117. 013	32.056	43. 004 40. 730 42. 125	1.00 20.05 1.00 13.15 1.00 16.85	В В В	0 C C
ATOM ATOM ATOM	9007 9008 9009	O N CA	THR TRP TRP	401 402 402	116.478	34. 519 36. 013 37. 199	43. 192 41. 659 42. 335	1.00 18.14 1.00 16.42 1.00 14.66	B B B	O N C
ATOM ATOM ATOM	9010 9011 9012		TRP TRP TRP	402 402 402	119.001		43. 561 43. 296 42. 861	1.00 16.17 1.00 16.85 1.00 17.78	B B B	C C C
ATOM ATOM ATOM	9013 9014 9015	CE3	TRP TRP TRP	402 402 402		38. 164 39. 948 36. 453	42. 771 42. 542 43. 440	1. 00 18. 27 1. 00 18. 13 1. 00 16. 20	B B B	C C C

ELC 4 105 (Continued										
				FIG. 4-185						
ATOM ATOM	9016 9017		402	100 100 00 000						
ATOM	9018		402 402	100 550						
ATOM	9019		402							
ATOM	9020		402							
ATOM	9021	0 TRP	402							
ATOM	9022		403	116.309 39.480 41.534 1.00 13.41 B						
ATOM ATOM	9023		403	116. 368 40. 554 40. 548 1. 00 12. 05 B	B C					
ATOM	9024 9025	CB GLU CG GLU	403 403	114 400 00 000						
ATOM	9026	CD GLU	403	110 000 00 000						
ATOM	9027	OE1 GLU	403	113. 288 39. 607 38. 391 1. 00 14. 00 B 112. 301 40. 306 38. 713 1. 00 15. 50 B						
ATOM	9028	OE2 GLU	403	113. 397 39. 068 37. 271 1. 00 14. 63 B						
ATOM	9029	C GLU	403	116. 852 41. 938 40. 999 1. 00 13. 29 B						
ATOM	9030	O GLU	403	116. 785 42. 301 42. 171 1. 00 14. 74 B						
ATOM ATOM	9031 9032	N VAL	404	117. 322 42. 716 40. 031 1. 00 12. 89 B	N					
ATOM	9033	CA VAL CB VAL	404 404	117. 800 44. 067 40. 270 1. 00 12. 91 B						
ATOM	9034	CG1 VAL	404	118. 926 44. 420 39. 265 1. 00 11. 91 B 119. 374 45. 859 39. 453 1. 00 13. 92 B						
ATOM	9035	CG2 VAL	404	119. 374 45. 859 39. 453 1. 00 13. 92 B 120. 096 43. 484 39. 459 1. 00 8. 31 B	C					
ATOM	9036	C VAL	404	116. 607 44. 994 40. 039 1. 00 14. 23 B	Č					
ATOM	9037	0 VAL	404	116. 129 45. 105 38. 918 1. 00 16. 13 B	ŏ					
ATOM ATOM	9038	N ILE	405	116. 122 45. 653 41. 089 1. 00 13. 56 B	Ň					
ATOM	9039 9040	CA ILE CB ILE	405 405	114. 968 46. 540 40. 951 1. 00 12. 56 B	C					
ATOM	9041	CG2 ILE	405	114. 453 47. 020 42. 339 1. 00 12. 98 B 113. 151 47. 763 42. 183 1. 00 7. 46 B	C					
ATOM	9042	CG1 ILE	405	113. 151 47. 763 42. 183 1. 00 7. 46 B 114. 256 45. 824 43. 282 1. 00 14. 03 B	C					
ATOM	9043	CD1 ILE	405		C C					
ATOM	9044	C ILE	405	115. 293 47. 762 40. 088 1. 00 14. 39 B	C					
ATOM	9045	0 ILE	405	114.504 48.156 39.226 1.00 14.58 B	ŏ					
ATOM ATOM	9046 9047	N GLY CA GLY	406	116. 455 48. 367 40. 315 1. 00 14. 30 B	N					
ATOM	9048	CA GLY C GLY	406 406	116. 822 49. 521 39. 521 1. 00 12. 80 B 118. 253 49. 967 39. 708 1. 00 13. 75 B	С					
ATOM	9049	0 GLY	406	110 0=0	C					
ATOM		N ILE	407	110 000 50 010 00 001	0					
ATOM	9051	CA ILE	407	118. 806 50. 618 38. 691 1. 00 14. 84 B 120. 161 51. 144 38. 760 1. 00 13. 37 B	N C					
ATOM	9052	CB ILE	407	120. 797 51. 192 37. 361 1. 00 11. 30 B	Č					
ATOM	9053	CG2 ILE	407	122. 039 52. 077 37. 373 1. 00 11. 29 B	Č					
ATOM ATOM	9054 9055	CG1 ILE CD1 ILE	407	121.163 49.768 36.936 1.00 9.82 B	C					
ATOM	9056	CDI ILE	407 407	121. 237 49. 545 35. 446 1. 00 9. 37 B 119. 991 52. 546 39. 343 1. 00 15. 02 B	C C					
ATOM	9057	0 ILE	407	110 000 #0 001						
ATOM	9058	N GLU	408	119. 236 53. 361 38. 819 1. 00 14. 39 B 120. 692 52. 825 40. 431 1. 00 16. 63 B	O N					
ATOM	9059	CA GLU	408	120.552 54.105 41.105 1.00 18.23 B	C					
ATOM	9060	CB GLU	408	120. 373 53. 849 42. 601 1. 00 21. 53 B	Č ·					
ATOM ATOM	9061 9062	CG GLU CD GLU	408	119. 290 52. 815 42. 906 1. 00 23. 80 B	C					
ATOM	9063	CD GLU OE1 GLU	408 408	117. 916 53. 275 42. 456 1. 00 27. 87 B	C					
ATOM	9064	OE2 GLU	408	117. 135 52. 429 41. 967 1. 00 30. 29 B 117. 612 54. 483 42. 598 1. 00 29. 06 B	0					
	-		-00	SUBSTITUTE SHEET (RULE 26)	0					
				,						

ATOM 9065 C GLU 408 121.687 55.094 40.888 1.00 19.22 B C ATOM 9066 O GLU 408 121.468 56.306 40.924 1.00 21.06 B O ATOM 9067 N ALA 409 122.899 54.589 40.678 1.00 18.36 B N ATOM 9068 CA ALA 409 124.698 54.583 40.473 1.00 18.36 B N ATOM 9068 CA ALA 409 124.533 56.012 41.816 1.00 16.78 B C ATOM 9070 C ALA 409 125.188 54.756 39.755 1.00 17.745 B C ATOM 9071 O ALA 409 125.323 55.366 39.755 1.00 17.745 B C ATOM 9071 O ALA 409 125.323 55.366 39.834 1.00 15.91 B O ATOM 9072 C LEU 410 126.009 55.545 39.662 1.00 17.35 B N ATOM 9073 CA LEU 410 127.140 55.034 38.311 1.00 17.53 B C ATOM 9073 CA LEU 410 127.140 55.034 38.311 1.00 17.53 B C ATOM 9075 CD LEU 410 127.767 54.292 38.862 1.00 18.12 B C ATOM 9076 CD LEU 410 127.767 54.292 38.862 1.00 18.12 B C ATOM 9076 CD LEU 410 127.144 54.224 34.467 1.00 16.50 B C ATOM 9077 CD LEU 410 128.2878 52.914 36.302 1.00 18.72 B C ATOM 9077 CD LEU 410 128.285 57.175 38.190 1.00 20.28 B C ATOM 9078 C LEU 410 128.285 57.175 38.190 1.00 20.28 B C ATOM 9078 C LEU 410 128.285 57.175 38.190 1.00 20.28 B C ATOM 9078 C LEU 410 128.2828 57.175 38.190 1.00 20.28 B C ATOM 9080 C LEU 410 128.2828 57.175 38.190 1.00 20.28 B C ATOM 9080 C LEU 410 128.228 57.175 38.190 1.00 20.28 B C ATOM 9080 C LEU 410 128.528 55.396 38.589 1.00 18.37 B N ATOM 9081 CA THR 411 129.532 55.396 38.589 1.00 18.37 B N ATOM 9082 CB THR 411 131.360 55.264 40.514 1.00 17.72 B C ATOM 9088 CA THR 411 131.360 55.264 40.514 1.00 17.72 B C ATOM 9088 CA THR 411 131.744 55.293 37.784 1.00 20.67 B C ATOM 9088 CA THR 411 131.744 55.293 37.784 1.00 20.67 B C ATOM 9098 CB SER 412 133.912 54.988 36.753 1.00 21.07 B N ATOM 9098 CB SER 412 133.915 5.772 37.543 1.00 21.07 B N ATOM 9099 C SER 412 133.915 5.772 37.543 1.00 21.07 B C ATOM 9099 C SER 412 133.915 5.772 37.543 1.00 21.07 B N ATOM 9099 C SER 412 133.915 5.772 37.543 1.00 22.07 B C ATOM 9099 C SER 412 133.915 5.772 37.543 1.00 22.07 B C ATOM 9099 C SER 412 134.961 52.843 36.955 1.00 22.77 B C ATOM 9099 C C SER 413 133.445 51.575 40.899 1.00 18.45 B C ATOM 9099 C C SER 413 133.446 51.		٠.				FIC	G. 4-	186			(Continued)
ATOM 9066 O GLU 408 121.468 56.306 40.924 1.00 21.06 B O ATOM 9067 N ALA 409 122.899 54.589 40.678 1.00 18.36 B N ATOM 9068 CA ALA 409 124.533 56.012 41.816 1.00 16.78 B C ATOM 9070 C ALA 409 124.533 56.012 41.816 1.00 16.78 B C ATOM 9071 O ALA 409 125.189 54.766 39.755 1.00 17.45 B C ATOM 9071 O ALA 409 125.323 53.536 39.834 1.00 15.91 B O ATOM 9072 N LEU 410 126.09 55.545 39.062 1.00 17.35 B N ATOM 9073 CA LEU 410 127.140 55.034 38.311 1.00 17.55 B N ATOM 9073 CA LEU 410 127.140 55.034 38.311 1.00 17.53 B C ATOM 9076 CD LEU 410 127.767 54.292 35.861 1.00 18.12 B C ATOM 9076 CD LEU 410 127.767 54.292 35.861 1.00 18.12 B C ATOM 9077 CG LEU 410 127.144 54.224 34.467 1.00 14.82 B C ATOM 9078 C LEU 410 128.278 52.914 36.302 1.00 16.12 B C ATOM 9077 C LEU 410 128.285 55.969 38.356 1.00 18.72 B C ATOM 9078 C LEU 410 128.285 55.969 38.356 1.00 18.72 B C ATOM 9078 C LEU 410 128.285 55.969 38.356 1.00 18.72 B C ATOM 9078 C LEU 410 128.285 55.969 38.356 1.00 18.72 B C ATOM 9078 C LEU 410 128.282 57.175 38.190 1.00 20.28 B O ATOM 9080 N THR 411 130.786 56.142 38.617 1.00 19.27 B C ATOM 9081 CA THR 411 130.786 56.142 38.617 1.00 19.27 B C ATOM 9082 CB THR 411 131.360 56.286 40.660 1.00 18.85 B C ATOM 9085 CT THR 411 131.374 54.200 37.576 1.00 17.72 B C ATOM 9085 C THR 411 131.374 54.200 37.577 1.00 23.60 B C ATOM 9088 CA SER 412 133.91 56.60 88 36.95 1.00 18.37 B N ATOM 9085 C THR 411 131.374 54.200 37.577 1.00 23.60 B C ATOM 9090 C SER 412 134.961 55.287 36.365 1.00 18.37 B C ATOM 9090 C SER 412 134.961 55.287 36.365 1.00 23.360 B C ATOM 9090 C SER 412 134.961 55.284 36.695 1.00 23.36 B C ATOM 9090 C SER 412 134.961 55.284 36.995 1.00 23.36 B C ATOM 9090 C SER 412 134.961 55.284 36.995 1.00 23.13 B C ATOM 9090 C SER 412 134.961 55.284 36.995 1.00 23.13 B C ATOM 9090 C SER 412 134.961 55.284 36.995 1.00 23.13 B C ATOM 9090 C SER 412 134.961 55.284 36.995 1.00 23.15 B C ATOM 9090 C SER 413 133.346 55.776 39.697 40.548 1.00 22.07 B C ATOM 9090 C SER 413 133.346 55.776 39.697 40.548 1.00 22.07 B C ATOM 9090 C SER 413 133.346	ATOM	9065	С	GLU	408				1.00 19.22	В	С
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ATOM 9107 CD2 TYR 414 133.093 51.850 43.718 1.00 14.91 B C ATOM 9108 CE2 TYR 414 134.071 51.282 44.512 1.00 16.48 B C ATOM 9109 CZ TYR 414 134.066 49.921 44.733 1.00 16.25 B C ATOM 9110 OH TYR 414 135.030 49.369 45.541 1.00 19.68 B O ATOM 9111 C TYR 414 129.787 51.898 40.214 1.00 17.91 B C ATOM 9112 0 TYR 414 129.547 52.990 39.693 1.00 17.06 B O										В	С
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ATOM 9109 CZ TYR 414 134.066 49.921 44.733 1.00 16.25 B C ATOM 9110 OH TYR 414 135.030 49.369 45.541 1.00 19.68 B O ATOM 9111 C TYR 414 129.787 51.898 40.214 1.00 17.91 B C ATOM 9112 O TYR 414 129.547 52.990 39.693 1.00 17.06 B O											
ATOM 9110 OH TYR 414 135.030 49.369 45.541 1.00 19.68 B O ATOM 9111 C TYR 414 129.787 51.898 40.214 1.00 17.91 B C ATOM 9112 O TYR 414 129.547 52.990 39.693 1.00 17.06 B O											
ATOM 9111 C TYR 414 129.787 51.898 40.214 1.00 17.91 B C ATOM 9112 0 TYR 414 129.547 52.990 39.693 1.00 17.06 B 0											
ATOM 9112 0 TYR 414 129.547 52.990 39.693 1.00 17.06 B 0											
Amore and a second seco											
ATOM 9113 N LEU 415 128.901 50.917 40.323 1.00 16.46 B N	ATOM	9113	N	LEU	415	128. 901					

	FIG. 4-187	(Continued)
ATOM 9114 CA LEU ATOM 9115 CB LEU ATOM 9116 CG LEU ATOM 9117 CD1 LEU ATOM 9118 CD2 LEU ATOM 9119 C LEU ATOM 9120 O LEU ATOM 9121 N TYR ATOM 9123 CB TYR ATOM 9124 CG TYR ATOM 9125 CD1 TYR ATOM 9126 CE1 TYR ATOM 9127 CD2 TYR ATOM 9128 CE2 TYR ATOM 9129 CZ TYR ATOM 9129 CZ TYR ATOM 9130 OH TYR ATOM 9131 C TYR ATOM 9131 C TYR ATOM 9132 O TYR ATOM 9133 N TYR ATOM 9134 CA TYR ATOM 9135 CB TYR ATOM 9136 CG TYR ATOM 9136 CG TYR ATOM 9137 CD1 TYR ATOM 9138 CE1 TYR ATOM 9138 CE1 TYR ATOM 9138 CE1 TYR ATOM 9139 CD2 TYR ATOM 9130 CD2 TYR ATOM 9130 CD2 TYR ATOM 9131 C TYR ATOM 9131 C TYR ATOM 9134 CA TYR ATOM 9135 CB TYR ATOM 9136 CG TYR ATOM 9137 CD1 TYR ATOM 9138 CE1 TYR ATOM 9138 CE1 TYR ATOM 9140 CE2 TYR ATOM 9140 CE2 TYR ATOM 9141 CZ TYR ATOM 9142 OH TYR ATOM 9143 C TYR ATOM 9144 O TYR ATOM 9145 N LLE ATOM 9146 CA LLE ATOM 9147 CB LLE ATOM 9148 CG2 LLE ATOM 9149 CG1 LLE ATOM 9140 CB1 LLE ATOM 9141 CB LLE ATOM 9140 CB1 LLE ATOM 9150 CD1 LLE	415 127.537 51.027 39.855 1.00 14.70 415 127.297 50.040 38.714 1.00 13.43 415 125.924 50.107 38.049 1.00 15.02 415 126.044 49.620 36.619 1.00 15.33 415 124.899 49.295 38.852 1.00 15.41 415 126.674 50.668 41.066 1.00 15.33 415 126.777 49.566 41.601 1.00 16.82 416 125.840 51.595 41.519 1.00 15.16 416 124.879 52.530 43.566 1.00 14.80 416 124.879 52.530 43.566 1.00 14.93 416 127.031 53.835 43.350 1.00 14.05 416 127.823 53.094 45.366 1.00 14.05 416 128.625 53.938 45.147 1.00 15.00 416 128.625 53.938 45.147 1.00 15.00 416 129.766 54.466 45.699 1.00 14.00 416 129.766 54.466 </td <td>Continued B C B C C B C C B C C B C C B C C B C C B C C B C C B C C B C C B C C B C C C B C</td>	Continued B C B C C B C C B C C B C C B C C B C C B C C B C C B C C B C C B C C B C C C B C
ATOM 9160 CA ASN ATOM 9161 CB ASN ATOM 9162 CG ASN	420 115.913 41.716 46.395 1.00 16.73 E 420 114.448 41.406 46.067 1.00 13.22 E 420 114.279 40.740 44.724 1.00 13.67 E	ВС

					5.5	-				(Continued)
					F I (G. 4	- 188	i		
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9163 9164 9165 9166 9167 9168 9169 9170 9171 9172 9173 9174 9175 9176	ND: C O N CA CB CC CD	1 ASN 2 ASN ASN GLU GLU GLU GLU GLU GLU GLU GLU GLU TYR TYR	420 420 421 421 421 421 421 421 421 421 421 421	115. 220 113. 072 116. 700 116. 135 118. 018 118. 895 120. 291 121. 358 122. 661 123. 169 123. 184 119. 028 118. 960 119. 223	40. 146 40. 818 40. 426 39. 368 40. 532 39. 393 39. 694 38. 747 38. 782 39. 890 37. 689 37. 756 39. 897	44. 193 44. 169 46. 638 46. 910 46. 543 46. 754 46. 734 45. 951 45. 661 45. 639 48. 218 48. 519 49. 120	1. 00 14. 68 1. 00 7. 98 1. 00 16. 85 1. 00 18. 37 1. 00 17. 15 1. 00 19. 34 1. 00 18. 78 1. 00 20. 75 1. 00 22. 48 1. 00 21. 18 1. 00 22. 04 1. 00 19. 80 1. 00 20. 89 1. 00 19. 38	B B B B B B B B B B B B B B B B B B B	0 N C O N C C C C C O
ATOM	9178	CB	TYR	$\begin{array}{c} 422 \\ 422 \end{array}$	119. 401 119. 386	39. 596 40. 895	50. 530 51. 326	1.00 19.16 1.00 19.06	B B	C C
ATOM ATOM ATOM	9179 9180 9181	CG CD1	TYR TYR TYR	422 422 422	119.881 121.046 121.510	40. 746 40. 023 39. 893	52. 741 53. 024 54. 314	1. 00 13. 60 1. 00 21. 59 1. 00 19. 84 1. 00 19. 73	B B B	C C C
ATOM	9182	CD2	2 TYR	422	119.198	41.334	53.798	1.00 21.32	В	C
ATOM ATOM	9183 9184	CZ	TYR TYR	$\begin{array}{c} 422 \\ 422 \end{array}$	119.658 120.813	41. 210 40. 488	55. 097 55. 347	1.00 23.82 1.00 23.64	B B	C C
ATOM	9185	ОН	TYR	422	121.267	40.376	56.637	1.00 28.92	В	0
ATOM ATOM	9186	C	TYR	422	118.401	38. 600	51.114	1.00 20.84	· B	C
ATOM	9187 9188	O N	TYR LYS	$\begin{array}{c} 422 \\ 423 \end{array}$	117. 187 118. 933	38. 779 37. 546	51. 012 51. 732	1.00 22.40 1.00 21.52	B	0
ATOM ·	9189	CA	LYS	423	118. 130	36.486	52. 340	1.00 21.52	B B	N C
ATOM	9190	CB	LYS	423	117.436	36. 995	53.608	1.00 22.83	В	Č
ATOM	9191	CG	LYS	423	118.393	37.278	54.751	1.00 25.85	B	č
ATOM	9192	CD	LYS	423	117.677	37.707	56.020	1.00 27.71	В	C
ATOM ATOM	9193 9194	CE NZ	LYS LYS	423 423	118.692	38.082	57.098	1.00 31.46	В	C .
ATOM	9195	C	LYS	423	118.052 117.097	38. 548 35. 906	58. 367 51. 378	1.00 31.96 1.00 21.44	В	N
ATOM	9196	Ŏ	LYS	423	116.114	35. 293	51. 797	1.00 21.44	В В	0 C
ATOM	9197	N	GLY	424	117. 331	36. 106	50.086	1.00 20.50	В	N N
ATOM	9198	CA	GLY	424	116.430	35.595	49.070	1.00 20.06	B	C
ATOM	9199	Ç	GLY	424	114.969	35.945	49. 274	1.00 20.45	В	Č
ATOM	9200	0	GLY	424	114.102	35. 120	49.013	1.00 21.91	В	0
ATOM ATOM	9201	N	MET	425	114.695	37. 163	49. 739	1.00 20.34	В	N
ATOM	9202 9203	CA CB	MET	425	113.322	37.627	49. 968	1.00 18.53	В	C
ATOM	9204	CG	MET MET	425 425	113. 234 113. 756	38. 329	51.317	1.00 19.68	В	C
ATOM	9205	SD	MET	425	113. 756	37.501	52. 469	1.00 22.38	В	C
ATOM	9206	CE	MET	425	111.741	38. 352 38. 663	54. 020 53. 907	1.00 24.27 1.00 21.26	В	S
ATOM	9207	C	MET	425	112. 908	38.604	48. 871	1.00 21.20	B B	C C
ATOM	9208	0 .	MET	425	113.405	39. 725	48. 819	1.00 10.73	В	0
ATOM	9209	N	PRO	426	111.968	38. 206	47. 999	1.00 16.64	В	N
ATOM	9210	CD	PRO	426	111.173	36. 969	48.017	1.00 17.29	В	C.
ATOM	9211	CA	PRO	426	111.530	39.089	46.910	1.00 15.29	B	Č

				FIG. 4-189	(Continued)
ATTOM	0010	ממ' מס			
ATOM ATOM	9212 9213			Б	C
ATOM	9214				C C
ATOM	9215			110. 913 41. 402 46. 727 1. 00 15. 90 B	ŏ
ATOM	9216			110. 362 40. 321 48. 630 1. 00 14. 46 B	N
ATOM ATOM	9217 9218			110 010	C
ATOM	9219			140 101 I	C
ATOM	9220				0 N
ATOM	9221	- CA GLY	428	112. 902 43. 036 50. 577 1. 00 8. 65 B	C
ATOM	9222			113. 735 43. 771 49. 538 1. 00 10. 35 B	č
ATOM ATOM	9223 9224			113. 778 43. 363 48. 377 1. 00 10. 03 B	0
ATOM	9225	N ARG CA ARG		114. 406 44. 844 49. 946 1. 00 11. 09 B 115. 224 45. 630 49. 023 1. 00 12. 98 B	N
ATOM	9226	CB ARG		115. 224 45. 630 49. 023 1. 00 12. 98 B 114. 349 46. 667 48. 314 1. 00 14. 68 B	C.
ATOM	-9227	CG ARG		113.580 46.084 47.144 1.00 18.95 B	C C
ATOM	9228	CD ARG		112. 423 46. 947 46. 701 1. 00 18. 69 B	Č
ATOM ATOM	9229	NE ARG		111.590 46.279 45.699 1.00 19.88 B	Ň
ATOM	9230 9231	CZ ARG NH1 ARG		111.184 45.008 45.769 1.00 21.09 B	C .
ATOM	9232	NH2 ARG		111. 535 44. 227 46. 791 1. 00 17. 36 B 110. 390 44. 520 44. 825 1. 00 20. 65 B	N
ATOM	9233	C ARG		110. 390 44. 520 44. 825 1. 00 20. 65 B 116. 420 46. 328 49. 678 1. 00 13. 64 B	N C
ATOM	9234	0 ARG	429	116. 291 46. 983 50. 707 1. 00 13. 96 B	0
ATOM	9235	N ASN	430	117. 584 46. 198 49. 056 1. 00 12. 81 B	Ň
ATOM ATOM	9236 9237	CA ASN CB ASN	430	118. 784 46. 812 49. 585 1. 00 13. 48 B	C
ATOM	9238	CG ASN	430 430	119. 605 45. 767 50. 344 1. 00 11. 94 B 118. 985 45. 411 51. 677 1. 00 12. 47 B	C
ATOM	9239	OD1 ASN	430	118. 985 45. 411 51. 677 1. 00 12. 47 B 119. 104 46. 167 52. 652 1. 00 11. 56 B	C
ATOM	9240	ND2 ASN	430	118. 293 44. 277 51. 727 1. 00 7. 39 B	O N
ATOM	9241	C ASN		119.644 47.477 48.528 1.00 14.50 B	Č
ATOM ATOM	9242 9243	O ASN	430	119. 530 47. 189 47. 335 1. 00 14. 26 B	0
ATOM	9244	N LEU CA LEU	431 431	120. 504 48. 377 48. 992 1. 00 16. 18 B	N
ATOM	9245	CB LEU	431	101 700 50 400 40 740	C
ATOM	9246		431	121. 709 50. 496 48. 713 1. 00 16. 67 B 122. 825 51. 279 48. 012 1. 00 18. 10 B	C C
ATOM	9247	CD1 LEU	431	122. 501 51. 399 46. 528 1. 00 17. 30 B	Č
ATOM	9248	CD2 LEU	431	122. 998 52. 651 48. 667 1. 00 14. 93 B	Č
ATOM ATOM	9249 9250	C LEU	431	122. 729 48. 338 48. 022 1. 00 17. 39 B	C
ATOM	9251	N TYR	431 432	123. 367 48. 018 49. 028 1. 00 19. 06 B 123. 112 48. 038 46. 789 1. 00 17. 62 B	0
ATOM	9252	CA TYR	432	123. 112 48. 038 46. 789 1. 00 17. 62 B 124. 344 47. 317 46. 511 1. 00 18. 05 B	N C
ATOM	9253	CB TYR	432	124. 061 45. 978 45. 826 1. 00 17. 24 B	C
ATOM	9254	CG TYR	432	123. 334 44. 944 46. 654 1. 00 18. 80 B	Č
ATOM ATOM	9255 9256	CD1 TYR CE1 TYR	432	121. 962 45. 034 46. 883 1. 00 19. 62 B	C
ATOM	9257	CD2 TYR	432 432	121. 289 44. 049 47. 601 1. 00 19. 23 B 124. 015 43. 843 47. 169 1. 00 17. 63 B	C
ATOM	9258	CE2 TYR	432	124.015 43.843 47.169 1.00 17.63 B 123.360 42.862 47.882 1.00 18.49 B	C
ATOM	9259	CZ TYR	432	121. 996 42. 968 48. 099 1. 00 20. 13 B	C C
ATOM	9260	OH TYR	432	121. 358 41. 994 48. 834 1. 00 21. 75 B	0
				SUBSTITUTE SHEET (RULE 26)	

					FIC	G. 4-	190			(Continued)
ATOM ATOM ATOM ATOM	9261 9262 9263 9264 9265 9266 9267 9270 9271 9272 9273 9274 9275 9276 9277 9278 9279 9280 9281 9282 9283 9284 9285 9286 9287 9288 9290 9291 9292 9293 9296	CG1 CD1 C O N CA CB CG CD OE1	ILE ILE ILE GLN GLN GLN GLN GLN GLN GLN LEU LEU LEU LEU LEU	432 433 433 433 433 433 433 433 433 433	F I (125. 193 124. 700 126. 474 127. 386 128. 237 129. 297 130. 239 131. 190 132. 101 128. 654 128. 654 129. 411 128. 645 128. 054 129. 580 128. 978 130. 646 130. 554 131. 804 133. 045 134. 253 135. 490 136. 715 136. 763 137. 713 133. 068 132. 969 133. 200 133. 197 133. 050 131. 785 131. 785 131. 785 131. 785 131. 748 130. 572 134. 391 134. 294	48. 142 49. 066 47. 805 48. 460 49. 536 49. 022 50. 146 49. 723 50. 834 47. 364 47. 364 46. 331 46. 429 44. 458 43. 676 46. 907 46. 264 47. 145 46. 220 46. 617 46. 429 47. 145 46. 154 46. 220 46. 617 47. 668 47. 527 48. 905 48. 90	45. 557 44. 903 45. 486 44. 563 45. 251 46. 606 47. 712 48. 104 44. 058 44. 820 42. 767 42. 191 41. 124 40. 061 40. 518 39. 379 41. 573 40. 915 41. 809	1. 00 17. 78 1. 00 18. 57 1. 00 16. 13 1. 00 14. 57 1. 00 16. 46 1. 00 16. 51 1. 00 16. 69 1. 00 17. 27 1. 00 13. 68 1. 00 11. 44 1. 00 13. 85 1. 00 15. 56 1. 00 14. 45 1. 00 14. 45 1. 00 14. 14 1. 00 14. 14 1. 00 14. 14 1. 00 14. 14 1. 00 16. 13 1. 00 17. 71 1. 00 18. 33 1. 00 20. 88 1. 00 21. 76 1. 00 24. 28 1. 00 25. 69 1. 00 26. 08 1. 00 26. 08 1. 00 27. 46 1. 00 19. 80 1. 00 19. 31 1. 00 18. 85 1. 00 25. 55 1. 00 27. 46	B B B B B B B B B B B B B B B B B B B	(Continued) C O N C C C C C C C C C C C C C C C C
ATOM ATOM	9297 9298 9299	N CA CB	SER SER SER	437 437 437	135. 517 136. 690 137. 967	46. 775 46. 069 46. 683	37. 613 37. 119 37. 689	1.00 26.98 1.00 26.89 1.00 26.26	B B B	N C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9300 9301 9302 9303 9304 9305 9306 9307 9308	OG C O N CA CB CG OD1 OD2 C	SER SER SER ASP ASP ASP ASP	437 437 437 438 438 438 438 438 438 438 438	137. 940 136. 593 137. 152 135. 882 135. 704 136. 702 136. 622 135. 517 137. 659 134. 286	46. 694 44. 597 43. 736 44. 310 42. 930 42. 588 41. 135 40. 557 40. 575 42. 691	39. 102 37. 507 36. 832 38. 595 39. 049 40. 151 40. 571 40. 495 40. 990 39. 572	1.00 31.19 1.00 27.29 1.00 29.17 1.00 26.66 1.00 26.32 1.00 28.65 1.00 30.81 1.00 32.19 1.00 33.46 1.00 24.90	B B B B B B B	O C C C C C C C C C C C C C C C C C C C

					FI	G 4	- 1 0 1			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9310 9311 9312 9313 9314 9315 9316 9317 9318 9319 9320	N CA CB CC CD . CE	TYR	439 439 439 439 439 439 439 439	F I 133. 959 133. 461 132. 083 131. 301 131. 420 131. 442 131. 322 131. 348 131. 405 131. 410	43. 060 42. 046 41. 780 41. 243 42. 125 43. 514 44. 329 41. 572 42. 379	38. 753 39. 123 37. 924 36. 698 36. 814 35. 687 35. 416 34. 285 34. 430		B B B B B B B	(Continued) O N C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9321 9322 9323 9324 9325 9326 9327 9328 9329 9330 9331 9332	C O N CA CB OG1	TYR TYR THR THR THR THR THR THR THR LYS LYS LYS	439 439 440 440 440 440 440 440 441 441 441	131. 928 130. 882 132. 953 132. 858 134. 102 135. 221 134. 418 132. 712 132. 169 133. 200 133. 123 134. 396	40. 823 40. 801 40. 030 39. 094 38. 196 38. 975 37. 568 39. 852 39. 328 41. 087 41. 905	40. 294 40. 933 40. 584 41. 699 41. 806 42. 250 40. 462 43. 014 43. 987 43. 039 44. 243	1.00 24.38 1.00 25.27 1.00 24.21 1.00 23.35 1.00 23.70 1.00 22.70 1.00 23.82 1.00 22.79 1.00 21.81 1.00 22.86 1.00 22.90	B B B B B B B B B B B	C O N C C O C C O N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9333 9334 9335 9336 9337 9338 9339 9340 9341 9342	CG CD CE NZ C O N CA CB CG1	LYS LYS LYS LYS LYS VAL VAL VAL	441 441 441 441 441 442 442 442 442	135. 620 136. 871 138. 053 139. 319 131. 881 131. 828 130. 880 129. 624 128. 458 127. 123	42. 741 41. 878 42. 702 41. 804 42. 577 42. 794 43. 891 42. 289 42. 984 42. 093 42. 770	44. 375 44. 682 44. 878 45. 201 45. 346 44. 329 43. 768 45. 039 45. 242 44. 799 45. 119	1.00 25.86 1.00 30.20 1.00 34.36 1.00 37.32 1.00 40.04 1.00 21.89 1.00 21.84 1.00 19.62 1.00 17.69 1.00 17.33 1.00 15.79	B B B B B B B B B B B B B B B B B B B	C C C N C O N C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9343 9344 9345 9346 9347 9348 9349 9350 9351 9352	C O N CA CB OG1 CG2 C	VAL VAL THR THR THR THR THR THR	442 442 443 443 443 443 443 443	128. 586 129. 502 129. 742 129. 129 129. 015 130. 040 131. 370 129. 923 127. 641 127. 210	41. 792 43. 299 42. 437 44. 528 44. 927 46. 035 45. 566 46. 442 45. 475 46. 483	43. 306 46. 733 47. 572	1. 00 11. 20 1. 00 20. 40 1. 00 22. 84 1. 00 20. 64 1. 00 22. 17 1. 00 24. 13 1. 00 28. 90 1. 00 22. 91 1. 00 23. 06 1. 00 26. 29	B B B B B B B	C C O N C C C C
ATOM ATOM ATOM ATOM ATOM ATOM	9353 9354 9355 9356 9357 9358	N CA C O CB SG	CYS CYS CYS CYS CYS	444 444 444 444 444 51	126. 948 125. 656 125. 963 126. 866 124. 801 123. 137 UBSTITUTE	44. 835 45. 368 46. 516 46. 411 44. 328 44. 986 SHEET	49. 754 50. 163 51. 115 51. 941 50. 878 51. 221 (RULE 26	1.00 21.88 1.00 22.22 1.00 20.79 1.00 19.89 1.00 24.50 1.00 27.42	B B B B	N C C O C S

			FIG. 4-192	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9360 CA LEG 9361 CB LEG 9362 CG LEG 9363 CD1 LEG 9364 CD2 LEG 9365 C LEG 9366 O LEG 9367 N SER 9368 CA SER 9369 CB SER 9370 OG SER 9371 C SER 9372 O SER 9373 N CYS 9374 CA CYS 9375 C CYS	445 445 445 445 445 446 446 446 446 446	125. 205 47. 602 51. 005 1. 00 20. 20 B 125. 442 48. 785 51. 824 1. 00 17. 71 B 125. 651 49. 988 50. 899 1. 00 15. 76 B 126. 714 49. 756 49. 812 1. 00 15. 86 B 126. 930 51. 008 48. 970 1. 00 13. 93 B 128. 007 49. 333 50. 480 1. 00 12. 34 B 124. 333 49. 099 52. 814 1. 00 19. 64 B 124. 446 50. 036 53. 608 1. 00 20. 41 B 123. 262 48. 314 52. 776 1. 00 21. 11 B 122. 131 48. 552 53. 656 1. 00 20. 24 B 120. 947 49. 077 52. 834 1. 00 20. 38 B 120. 577 48. 143 51. 829 1. 00 18. 25 B 121. 708 47. 307 54. 411 1. 00 20. 86 B 121. 085 47. 404 55. 463 1. 00 21. 91 B 122. 043 46. 141 53. 874 1. 00 23. 05 B 121.	N C C C O N C C O N C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9376 O CYS 9377 CB CYS 9378 SG CYS 9379 N GLU 9380 CA GLU 9381 CB GLU 9382 CG GLU 9383 CD GLU 9384 OE1 GLU 9385 OE2 GLU 9386 C GLU	447 447 448 448 448 448 448 448 448 448	120. 881 44. 602 56. 739 1. 00 24. 50 B 122. 461 43. 722 53. 874 1. 00 24. 68 B 122. 134 43. 458 52. 103 1. 00 31. 64 B 123. 080 45. 011 56. 463 1. 00 23. 42 B 123. 394 44. 913 57. 881 1. 00 23. 49 B 124. 805 44. 358 58. 061 1. 00 24. 37 B 125. 060 43. 017 57. 395 1. 00 28. 24 B 123. 996 41. 985 57. 713 1. 00 34. 11 B 123. 377 42. 073 58. 796 1. 00 36. 48 B 123. 786 41. 070 56. 882 1. 00 37. 27 B	0 C S N C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9387 O GLU 9388 N LEU 9389 CA LEU 9390 CB LEU 9391 CG LEU 9392 CD1 LEU 9393 CD2 LEU 9394 C LEU 9395 O LEU 9396 N ASN	448 449 449 449 449 449 449 449 450	123. 249 46. 162 58. 738 1. 00 23. 12 B 123. 458 46. 101 59. 948 1. 00 24. 21 B 122. 900 47. 289 58. 134 1. 00 20. 81 B 122. 733 48. 516 58. 899 1. 00 20. 59 B 122. 123 49. 592 58. 010 1. 00 18. 76 B 123. 019 50. 143 56. 909 1. 00 17. 31 B 122. 221 51. 089 56. 045 1. 00 18. 95 B 124. 199 50. 868 57. 527 1. 00 16. 25 B 121. 853 48. 311 60. 144 1. 00 22. 20 B 122. 232 48. 674 61. 261 1. 00 22. 97 B 120. 677 47. 731 59. 937 1. 00 22. 75 B	C O N C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9397 CA ASN 9398 CB ASN 9399 CG ASN 9400 OD1 ASN 9401 ND2 ASN 9402 C ASN 9403 O ASN 9404 N PRO 9405 CD PRO 9406 CA PRO 9407 CB PRO	450 450 450 450 450 450 451 451 451	119. 729 47. 462 61.011 1.00 21.80 B 118. 958 48. 731 61.344 1.00 23. 73 B 118. 226 48. 632 62. 661 1.00 26. 67 B 117. 678 47. 581 63. 004 1.00 26. 78 B 118. 199 49. 733 63. 406 1.00 26. 73 B 118. 772 46. 400 60. 469 1.00 22. 01 B 117. 649 46. 701 60. 072 1.00 21. 48 B 119. 215 45. 134 60. 442 1.00 21. 65 B 120. 506 44. 673 60. 969 1.00 20. 73 B 118. 430 44. 004 59. 941 1.00 21. 39 B 119. 362 42. 817 60. 162 1.00 19. 94 B	C C C O N C O N C C C

ATOM 9408 CG PRO 451 120.209 43.253 61.290 1.00 21.78 B C ATOM 9409 C PRO 451 117.035 43.774 60.509 1.00 23.49 B C ATOM 9410 N PRO 451 116.125 43.372 59.774 1.00 25.66 B O ATOM 9411 N GLU 452 116.830 44.003 61.800 1.00 24.25 B N ATOM 9412 CA GLU 452 115.539 43.793 61.800 1.00 24.25 B N ATOM 9412 CA GLU 452 116.661 42.720 64.455 10.00 33.54 B C ATOM 9414 CC GLU 452 116.661 42.720 64.455 10.00 33.54 B C ATOM 9416 OEI GLU 452 116.661 42.720 64.455 10.00 33.54 B C ATOM 9416 OEI GLU 452 116.661 42.720 66.521 1.00 44.38 B C ATOM 9417 082 GLU 452 116.610 94.85 116.010 43.85 B C ATOM 9418 C GLU 452 116.610 94.85 116.010 43.85 B C ATOM 9419 0 GLU 452 114.543 44.867 61.968 1.00 25.59 B C ATOM 9419 0 GLU 452 113.374 44.867 61.968 1.00 27.44 B O ATOM 9420 C A ARG 453 114.132 47.198 61.478 1.00 21.67 B C ATOM 9421 C A ARG 453 114.132 47.198 61.478 1.00 21.67 B C ATOM 9420 C A ARG 453 114.132 47.198 61.478 1.00 21.67 B C ATOM 9421 C A ARG 453 114.132 47.198 61.478 1.00 21.67 B C ATOM 9422 C A ARG 453 114.165 50.878 62.65.521 1.00 47.19 B C ATOM 9421 C A ARG 453 114.165 50.878 62.652 10.071.23 B C ATOM 9422 C A ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9422 C A ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9422 C A ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9422 C A ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9423 C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9425 C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9427 C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9428 C C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9428 C C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9428 C C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9428 C C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9427 C C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9428 C C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9428 C C ARG 453 114.65 50.878 62.652 10.071.23 B C ATOM 9428 C C ARG 453 114.65 60 88.65 10.071.23 B C ATOM 9428 C C ARG 453 114.65 60 88.65 10.071.23 B C ATOM 9428 C C ARG 453 114.65 60 88.65 10.071.23 B		•				
ATOM 9409 C PRO 451 117.035 43.774 60.509 1.00 23.49 B C ATOM 9410 N GLU 452 116.125 43.392 9.774 1.00 25.06 B O ATOM 9412 CA GLU 452 115.539 43.783 62.394 1.00 26.56 B C ATOM 9413 CB GLU 452 115.650 43.787 69.597 1.00 44.38 B C ATOM 9416 OBI GLU 452 115.650 43.787 69.597 1.00 44.38 B C ATOM 9416 OBI GLU 452 116.661 42.70 64.455 1.00 39.54 B C ATOM 9417 0E2 GLU 452 117.355 41.782 66.521 1.00 44.38 B C ATOM 9419 OBI GLU 452 116.619 43.829 66.627 1.00 44.38 B C ATOM 9419 OBI GLU 452 116.619 43.829 66.627 1.00 44.38 B C ATOM 9419 OBI GLU 452 116.619 43.829 66.627 1.00 46.89 B O ATOM 9419 OBI GLU 452 114.543 44.867 61.988 1.00 25.59 B C ATOM 9419 OBI GLU 452 114.543 44.867 61.988 1.00 25.59 B C ATOM 9420 N ARG 453 115.010 46.101 61.848 1.00 23.36 B N ATOM 9421 CA ARG 453 114.132 47.198 61.478 1.00 21.67 B C ATOM 9422 CB ARG 453 114.132 47.198 61.478 1.00 21.67 B C ATOM 9424 CD ARG 453 113.364 52.088 62.234 1.00 20.24 B C ATOM 9424 CD ARG 453 114.132 47.198 61.284 1.00 21.67 B C ATOM 9424 CD ARG 453 114.152 94.8463 62.234 1.00 20.24 B C ATOM 9424 CD ARG 453 114.152 94.198 62.297 1.00 17.23 B C ATOM 9424 CD ARG 453 113.582 53.245 62.297 1.00 17.23 B C ATOM 9424 CD ARG 453 114.165 50.878 62.297 1.00 17.23 B C ATOM 9420 N ARG 453 114.165 50.878 62.297 1.00 17.21 B C ATOM 9420 N ARG 453 114.165 50.878 62.297 1.00 17.21 B C ATOM 9420 C ARG 453 113.582 53.245 62.997 1.00 17.21 B C ATOM 9430 O ARG 453 114.077 47.527 59.994 1.00 21.78 B C ATOM 9430 O ARG 453 114.077 47.527 59.994 1.00 21.78 B C ATOM 9430 O ARG 453 114.077 47.527 59.994 1.00 12.18 B C ATOM 9430 O ARG 453 114.077 47.527 59.994 1.00 12.18 B C ATOM 9430 O ARG 453 114.077 47.527 59.994 1.00 12.18 B C ATOM 9430 O ARG 453 114.077 47.527 59.994 1.00 14.66 B N ATOM 9430 C CYS 454 115.296 47.388 59.312 1.00 19.47 B C ATOM 9430 O ARG 453 114.077 47.527 59.994 1.00 14.66 B N ATOM 9430 C CYS 454 115.296 47.388 59.312 1.00 19.47 B C ATOM 9440 C CYS 454 115.596 50.305 56.505 1.00 19.47 B C ATOM 9441 CD GLN 455 114.685 45.895 56.505 1.00 19.47 B C ATOM 9440 C CYS 454 115.6				FIG. 4-193	(Continu	ıed)
ATOM 9453 CE2 TYR 456 110.207 49.046 49.200 1.00 4.15 B C ATOM 9454 CZ TYR 456 109.669 47.910 48.629 1.00 8.20 B C ATOM 9455 OH TYR 456 108.949 47.994 47.464 1.00 11.71 B 0	ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9409 C PR 9410 O PR 9411 N GL 9412 CA GL 9413 CB GL 9414 CG GL 9415 CD GL 9416 OE1 GL 9417 OE2 GL 9418 C GL 9419 O GLI 9419 CA ARC 9421 CA ARC 9422 CB ARC 9423 CG ARC 9424 CD ARC 9425 NE ARC 9426 CZ ARC 9427 NH1 ARC 9428 NH2 ARC 9428 NH2 ARC 9429 C ARC 9430 O ARC 9431 N CYS 9432 CA CYS 9433 C CYS 9434 O CYS 9435 CB CYS 9436 SG CYS 9437 N GLN 9439 CB GLN 9439 CB GLN 9440 CG GLN 9441 CD GLN 9442 OE1 GLN 9443 NE2 GLN 9444 C GLN 9445 O GLN 9446 N TYR 9446 N TYR 9447 CA TYR 9448 CB TYR 9450 CD1 TYR 9451 CE1 TYR	451 452 452 452 452 452 453 453 453 453 455 455 455 455 455 455	120. 209 43. 253 61. 290 1. 00 21. 78 117. 035 43. 774 60. 509 1. 00 23. 49 116. 125 43. 392 59. 774 1. 00 25. 06 116. 850 44. 003 61. 800 1. 00 24. 25 115. 539 43. 793 62. 394 1. 00 26. 56 115. 650 43. 767 63. 920 1. 00 32. 21 116. 621 42. 720 64. 455 1. 00 39. 54 116. 616 42. 675 65. 976 1. 00 47. 19 116. 019 43. 529 66. 627 1. 00 47. 19 116. 019 43. 529 66. 627 1. 00 47. 19 116. 019 43. 529 66. 627 1. 00 47. 19 115. 010 46. 101 61. 848 1. 00 23. 36 114. 132 47. 198 61. 478 1. 00 21. 67 114. 539 48. 463 62. 234 1. 00 21. 94 113. 714 49. 685 61. 872 1. 00 20. 24 114. 165 50. 878 <td>C C C O N C C C C O O C O N C C C C C N C N</td> <td>ied)</td>	C C C O N C C C C O O C O N C C C C C N C N	ied)
1-0 101 00. 100 1 1. UT 1 1 U	ATOM ATOM ATOM	9453 CE2 TYR 9454 CZ TYR 9455 OH TYR	456 456 456	110. 207 49. 046 49. 200 1. 00 4. 15 109. 669 47. 910 48. 629 1. 00 8. 20	B C C	

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				٠.	FIC	G. 4-	194			(Continued)
ATOM ATOM	9457 9458	0 N	TYR TYR	456 457	113. 127 114. 752	49. 991 49. 309	52. 775 51. 382	1.00 15.30 1.00 15.11	В	0
ATOM	9459	CA	TYR	457	115. 286	50. 646	51. 152	1.00 15.11	В	N
ATOM	9460	CB	TYR	457	116. 792	50.674	51. 390	1.00 14.63	B B	C
ATOM	9461	CG	TYR	457	117. 271	50. 394	52. 786	1.00 14.51	В	C C C C C C
ATOM	9462	CD1		457	117. 364	49. 088	53. 275	1.00 14.02	В	C
ATOM	9463		TYR	457	117. 903	48. 836	54. 540	1.00 14.47	В	C
ATOM	9464		TYR	457	117.714	51.434	53. 595	1.00 13.34	В	C
ATOM	9465		TYR	457	118. 245	51. 193	54. 850	1.00 13.54	В	Č.
ATOM	9466	CZ	TYR	457	118.341	49. 902	55. 318	1.00 11.72	В	Č
ATOM	9467	OH	TYR	457	118.877	49. 701	56. 559	1.00 8.57	B	ő
ATOM	9468	Č	TYR	457	115.085	51. 192	49.742	1.00 15.66	В	č
ATOM	9469	0	TYR	457	114.827	50.455	48.797	1.00 17.46	B	Ö
ATOM	9470	N	SER	458	115.234	52.505	49.624	1.00 14.42	B	N
ATOM	9471	CA	SER	458	115.176	53. 207	48.352	1.00 14.00	В	C
ATOM	9472	CB	SER	458	113.853	53.950	48.163	1.00 12.81	В	C
ATOM	9473	0G	SER	458	113.804	55.138	48.932	1.00 15.84	В	0
ATOM	9474	C	SER	458	116.318	54. 175	48.620	1.00 15.10	В	C
ATOM	9475	0	SER	458	116.631	54. 431	49. 791	1.00 14.29	В	0
ATOM	9476	N	VAL	459	116.946	54. 709	47.574	1.00 13.45	В	Ν .
ATOM	9477	CA	VAL	459	118.086	55. 593	47.779	1.00 13.00	В	C
ATOM	9478	CB	VAL	459	119.392	54. 853	47. 433	1.00 13.28	В	C
ATOM	9479	CG1		459	119.442	54.578	45. 934	1.00 10.72	В	C
ATOM	9480		VAL	459	120.600	55.672	47.878	1.00 13.89	В	C
ATOM	9481	C	VAL	459	118.051	56.882	46.969	1.00 14.23	В	C
ATOM ATOM	9482 9483	O N	VAL SER	459 460	117. 283	57.007	46.021	1.00 14.51	В	0
ATOM	9484	CA	SER	460	118.901 118.997	57.834 59.106	47. 347	1.00 14.01	В	N
ATOM	9485	CB	SER	460	118.039	60.116	46. 643 47. 272	1.00 14.81 1.00 15.45	В	C
ATOM	9486	OG	SER	460	118.038	61.333	46. 553	1.00 13.45	B B	C
ATOM	9487	C	SER	460	120.442	59. 629	46.693	1.00 15.07	В	C C
ATOM	9488	ŏ	SER	460	120. 930	60.040	47. 752	1.00 13.13	В	0
ATOM	9489	Ň	PHE	461	121.123	59.611	45. 547	1.00 14.73	В	N N
ATOM	9490	CA	PHE	461	122.516	60.068	45. 469	1.00 14.06	B	Ç .
ATOM	9491	CB	PHE	461	123.314	59. 229	44. 454	1.00 10.57	В	č
ATOM	9492	CG	PHE	461	123.583	57.809	44. 885	1.00 8.39	B	č
ATOM	9493	CD1	PHE	461	122.594	56.832	44.792	1.00 7.71	B	č
ATOM	9494	CD2	PHE	461	124.837	57.444	45.367	1.00 6.73	B	č
ATOM	9495		PHE	461	122.848	55.509	45.172	1.00 6.28	В	Č
ATOM	9496		PHE	461	125.105	56.118	45.752	1.00 6.24	В	Ċ
ATOM	9497	CZ	PHE	461	124.108	55.153	45.653	1.00 6.94	В	С
ATOM	9498	C	PHE	461	122.665	61.533	45.066	1.00 16.79	В	C C C C C
ATOM	9499	0	PHE	461	121.833	62.076	44.340	1.00 17.81	В	0
ATOM	9500	N	SER	462	123.740	62.170	45.528	1.00 18.84	В	N
ATOM	9501	CA	SER	462	124. 019	63. 555	45. 155	1.00 20.51	В	C
ATOM	9502	CB	SER	462	125. 131	64. 137	46.036	1.00 21.92	В	C
ATOM	9503	OG C	SER	462	126. 346	63. 421	45.878	1.00 24.40	В	0
ATOM	9504	C	SER	462	124. 465	63. 559	43.687	1.00 20.69	В	C
ATOM	9505	0	SER	462	124.607	62. 505	43.075	1.00 21.27	В	0

					(Continued)
		•		FIG. 4-196	(3022322000,
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9555 9556 9556 9556 9556 9561 9562 9563 9563 9566 9566 9571 9577 9577 9577 9577 9578 9581 9583 9583 9583 9583 9583 9593 9593 9594	CD2 TYR CE2 TYR	468 468 468 468 469 469 469 469 469 470 470 470 470 470 471 471 471 471 471 471 471 471 471 471	120. 369 59. 235 51. 355 1. 00 18. 78 B 119. 277 60. 302 51. 283 1. 00 16. 79 B 118. 247 60. 143 52. 393 1. 00 16. 33 B 117. 035 61. 034 52. 214 1. 00 16. 44 B 116. 438 61. 076 51. 147 1. 00 18. 52 B 116. 659 61. 739 53. 265 1. 00 16. 60 B 119. 729 57. 855 51. 240 1. 00 18. 75 B 119. 353 57. 413 50. 156 1. 00 20. 25 B 119. 641 57. 160 52. 359 1. 00 18. 03 B 119. 871 54. 860 53. 153 1. 00 12. 88 B 120. 920 54. 116 52. 334 1. 00 7. 18 B 121. 669 53. 176 53. 230 1. 00 9. 83 B 120. 248 53. 344 51. 241 1. 00 5. 95 B 117. 573 56. 769 54. 082 1. 00 17. 50 B 116. 644 55. 437 52. 517 1. 00 20. 97 B 115. 306	
ATOM	9593	0 CYS	472	112. 732 52. 636 55. 321 1. 00 21. 14 B 112. 323 53, 547 56. 036 1. 00 21. 12 B	C 0
ATOM ATOM	9594 9595 9596	SG CYS	472	114. 981 52. 073 56. 299 1. 00 23. 91 B 114. 149 50. 907 57. 416 1. 00 27. 85 B	C S
ATOM ATOM	9597 9598	N SER CA SER CB SER	473 473 473	111. 919 51. 755 54. 756 1. 00 19. 44 B 110. 482 51. 846 54. 967 1. 00 18. 92 B 109. 789 52. 191 53. 646 1. 00 18. 36 B	'N C
ATOM ATOM	9599 9600	OG SER C SER	473 473	110. 141 51. 261 52. 642 1. 00 21. 93 B	C 0
ATOM ATOM	9601 9602	O SER N GLY	473 474	109. 832 50. 609 55. 581 1. 00 17. 21 B 108. 615 50. 465 55. 530 1. 00 19. 59 B 110. 629 49. 716 56. 156 1. 00 16. 48 B	C 0 N
ATOM	9603	CA GLY	474	110.055 48.532 56.771 1.00 16.90 B	N C

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					FIC	G. 4-	198			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9653 9654 9655 9656 9657 9658 9660 9661 9662 9663 9664 9665 9666 9667 9668 9669 9670 9671	C O N CA CB CCD C C O N CA CB CC CD C C C CD C C CD C CD CD CD CD CD	THR THR THR THR LEU LEU LEU LEU HIS HIS HIS HIS	481 481 481 481 482 482 482 482 482 482 482 483 483 483 483	120. 129 120. 774 120. 459 120. 256 120. 964 120. 650 122. 035 122. 937 123. 203 123. 765 124. 115 124. 243 124. 684 124. 849 126. 090 125. 791 124. 697 123. 358 124. 933	58. 431 57. 163 56. 065 56. 864 58. 919 58. 648 59. 646 60. 166 61. 653 62. 439 62. 475 63. 856 59. 373 59. 013 59. 096 58. 332 56. 894 56. 245 56. 434 55. 258	56. 924 57. 480 56. 622 58. 858 55. 752 54. 602 56. 058 55. 038 55. 279 54. 092 52. 975 54. 525 55. 121 56. 210 53. 970 53. 970 53. 93 53. 488 54. 276 54. 264 55. 211	1. 00 15. 65 1. 00 14. 54 1. 00 18. 10 1. 00 15. 87 1. 00 16. 24 1. 00 16. 93 1. 00 18. 90 1. 00 19. 21 1. 00 20. 10 1. 00 21. 90 1. 00 21. 10 1. 00 22. 66 1. 00 19. 39 1. 00 20. 79 1. 00 18. 33 1. 00 16. 79 1. 00 14. 55 1. 00 14. 89 1. 00 15. 13 1. 00 16. 09	B B B B B B B B B B B B B B B B B B B	Continued) C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM	9673 9674	CE1	HIS HIS	483 483 483	124. 933 123. 788 122. 816	55. 258 54. 867 55. 565	55. 211 55. 736 55. 178	1.00 16.09 1.00 13.84 1.00 14.31	В В В	N C N
ATOM ATOM ATOM	9675 9676 9677	C O N	HIS HIS SER	483 483 484	127. 043 126. 617 128. 333	58. 939 59. 665 58. 645	52.868 51.961 53.003	1. 00 18. 94 1. 00 19. 56 1. 00 19. 52	B B B	C · · O N
ATOM ATOM ATOM ATOM	9678 9679 9680 9681	CA CB OG C	SER SER SER SER	484 484 484 484	129. 318 130. 520 131. 351 129. 774	59. 131 59. 779 58. 803 57. 907	52. 040 52. 738 53. 344 51. 259	1.00 21.33 1.00 21.77 1.00 24.25 1.00 21.22	В В В В	C C O C
ATOM ATOM ATOM ATOM	9682 9683 9684 9685	O N CA CB	SER SER SER SER	484 485 485 485	129. 942 129. 979 130. 389 130. 095	56. 827 58. 076 56. 967 57. 301	51.830 49.960 49.110 47.645	1.00 19.26 1.00 22.12 1.00 25.62 1.00 26.28	B B B	O N C C
ATOM ATOM ATOM ATOM	9686 9687 9688 9689	OG C O N	SER SER SER VAL	485 485 485 486	128. 715 131. 840 132. 097 132. 781	57. 552 56. 495 55. 300 57. 416	47. 444 49. 221 49. 138	1.00 30.40 1.00 26.33 1.00 27.23	B B B	0 . C 0
ATOM ATOM ATOM	9690 9691 9692	CA CB CG1	VAL VAL VAL	486 486 486	134. 194 135. 084 134. 786	57. 056 58. 284 58. 797	49. 407 49. 468 49. 798 51. 192	1.00 28.07 1.00 29.41 1.00 30.37 1.00 31.49	B B B	N C C C
ATOM ATOM ATOM ATOM	9693 9694 9695 9696	CG2 C O N	VAL VAL VAL ASN	486 486 486 487	136. 553 134. 507 135. 269 133. 922	57. 909 55. 929 55. 016 55. 979	49. 665 50. 442 50. 119 51. 630	1. 00 30. 81 1. 00 30. 57 1. 00 31. 62 1. 00 30. 95	B B B	C C 0
ATOM ATOM ATOM	9697 9698 9699	CA CB CG	ASN ASN ASN	487 487 487	134. 159 134. 888 136. 336	54. 928 55. 498 55. 868	52. 610 53. 833 53. 537	1.00 30.35 1.00 31.75 1.00 35.87 1.00 38.55	В В В	N C C C
ATOM ATOM	9700 9701	OD1 ND2		487 487	136. 838 137. 019	56. 895 55. 026	54. 014 52. 759	1.00 38.47 1.00 37.49	B B	O N

Continued)											
					F I G. 4 - 199						
ATOM	9702		ASN	487	132. 850 54. 288 53. 048 1. 00 30. 74 B	· C					
ATOM ATOM	9703		ASN	487	132. 830 53. 486 53. 982 1. 00 31. 45 B	0					
ATOM	9704 9705		ASP	488	131. 762 54. 633 52. 364 1. 00 28. 68 B 130. 449 54. 108 52. 707 1. 00 26. 66 B	N					
ATOM	9706	CB		488 488	D	C					
ATOM	9707	CG		488	130. 331 52. 636 52. 313 1. 00 27. 90 B 130. 253 52. 440 50. 816 1. 00 29. 72 B	C					
ATOM	9708		1 ASP	488							
ATOM	9709		2 ASP	488	129. 461 53. 146 50. 161 1. 00 31. 30 B 130. 977 51. 572 50. 290 1. 00 32. 18 B	0					
ATOM	9710	C	ASP	488	130. 219 54. 259 54. 204 1. 00 25. 72 B	0 C					
ATOM	9711	0	ASP	488	129. 654 53. 382 54. 856 1. 00 24. 30 B	0					
ATOM	9712	N	LYS	489	130. 669 55. 378 54. 754 1. 00 25. 25 B	N N					
ATOM	9713	CA		489	130. 503 55. 610 56. 176 1. 00 24. 10 B	Ċ					
ATOM	9714	CB	LYS	489	131. 607 56. 529 56. 705 1. 00 24. 94 B	Č					
ATOM	9715	CG	LYS	489	131.622 57.898 56.069 1.00 29.19 B	Ċ					
ATOM	9716	CD	LYS	489	132. 805 58. 719 56. 560 1. 00 33. 11 B	C					
ATOM	9717	CE	LYS	489	132.771 60.133 55.995 1.00 34.94 B	. С					
ATOM ATOM	9718	NZ	LYS	489	133.883 60.959 56.541 1.00 39.70 B	N					
ATOM	9719 9720	C 0	LYS LYS	489 489	100	C					
ATOM	9721	N	GLY	490	128. 556 56. 872 55. 585 1. 00 20. 15 B 128. 639 55. 968 57. 657 1. 00 22. 04 B	0					
ATOM	9722	CA	GLY	490	40= 0=0	Ň					
ATOM	9723	C	GLY	490	127. 352 56. 487 58. 067 1. 00 20. 03 B 127. 545 57. 854 58. 676 1. 00 20. 18 B	C C					
ATOM	9724	0	GLY	490	128. 091 57. 989 59. 769 1. 00 20. 54 B	0					
ATOM	9725	N	LEU	491	127. 092 58. 876 57. 965 1. 00 19. 44 B	N N					
ATOM	9726	CA	LEU	491	127. 234 60. 233 58. 440 1. 00 19. 54 B	Č					
ATOM	9727	CB	LEU	491	127. 032 61. 203 57. 283 1. 00 20. 53 B	Č					
ATOM	9728	CG	LEU	491	128. 153 61. 167 56. 242 1. 00 18. 39 B	Č					
ATOM	9729		LEU	491	127. 831 62. 089 55. 090 1. 00 19. 23 B	Ċ					
ATOM ATOM	9730		LEU	491	129. 441 61. 577 56. 898 1. 00 18. 31 B	C					
ATOM	9731 9732	C 0	LEU LEU	491	126. 287 60. 555 59. 586 1. 00 20. 91 B	C					
ATOM	9733	N	ARG	491 492	126. 735 60. 780 60. 713 1. 00 22. 15 B	0					
ATOM	9734	CA	ARG	492	124. 984 60. 566 59. 316 1. 00 20. 73 B 124. 020 60. 881 60. 364 1. 00 20. 06 B	N					
ATOM	9735	CB	ARG	492	104 000 00 000	C					
ATOM	9736	CG	ARG	492	100 000 00 011 00 000	C					
ATOM	9737	CD	ARG	492	123. 393 63. 244 59. 568 1. 00 20. 08 B 123. 759 64. 698 59. 798 1. 00 21. 15 B	C C					
ATOM	9738	NE	ARG	492	125. 193 64. 888 59. 625 1. 00 21. 60 B	N N					
ATOM	9739	CZ	ARG	492	125. 765 65. 192 58. 466 1. 00 23. 12 B	C					
ATOM	9740	NH1	ARG	492	125. 022 65. 360 57. 380 1. 00 24. 47 B	Ň					
ATOM	9741		ARG	492	127. 083 65. 286 58. 383 1. 00 23. 72 B	N					
ATOM	9742	C	ARG	492	122. 585 60. 443 60. 085 1. 00 21. 47 B	Č					
ATOM	9743	0	ARG	492	122. 247 59. 998 58. 983 1. 00 21. 32 B	0					
ATOM	9744	N	VAL	493	121. 746 60. 580 61. 107 1. 00 20. 97 B	N					
ATOM ATOM	9745 9746	CA CB	VAL VAL	493 493	120.344 60.211 61.018 1.00 21.38 B	Č .					
ATOM	9747	.CG1		493 493	119. 883 59. 537 62. 325 1. 00 22. 41 B	C					
ATOM	9748		VAL	493	118. 402 59. 215 62. 247 1. 00 23. 17 B 120. 698 58. 266 62. 574 1. 00 20. 83 B	C					
ATOM	9749	C	VAL	493	20,00	C					
ATOM	9750	Ö	VAL	493	119. 497 61. 456 60. 763 1. 00 21. 55 B 119. 462 62. 371 61. 580 1. 00 21. 85 B	. O					
		-	_		SUBSTITUTE SHEET (RULE 26)	U					

			EIC 4 200	(Continued)
			FIG. 4-200	
ATOM ATOM ATOM ATOM	9752 CA LE 9753 CB LE	U 494 U 494	118.811 61.485 59.626 1.00 21.18 117.974 62.626 59.264 1.00 19.43 117.782 62.660 57.742 1.00 19.57 119.101 62.610 56.953 1.00 21.60	B N B C B C
ATOM	9755 CD1 LE	U 494	119.101 62.610 56.953 1.00 21.60 118.832 62.502 55.456 1.00 20.38	B C B C
ATOM ATOM			119. 929 63. 851 57. 271 1. 00 19. 91	B C
ATOM	9758 0 LE		116.615 62.576 59.964 1.00 18.58 116.111 63.595 60.443 1.00 18.81	B C B O
ATOM			116.025 61.390 60.022 1.00 16.24	B N
ATOM ATOM			114. 729 61. 225 60. 659 1. 00 16. 44	B C
ATOM			113.612 61.651 59.698 1.00 17.53 112.217 61.506 60.268 1.00 19.67	B C B C
ATOM	9763 CD GLI	J 495	111.984 62.399 61.476 1.00 22.97	B C C
ATOM ATOM	9764 OE1 GLI 9765 OE2 GLI		112.023 63.642 61.315 1.00 22.51	B 0
ATOM	9766 C GLI			B O B C
ATOM	9767 0 GLU	495	114.678 58.875 60.236 1.00 15.37	B C B 0
ATOM ATOM	9768 N ASF 9769 CA ASF		114. 264 59. 534 62. 329 1. 00 14. 29	B N
ATOM	9770 CB ASP		11E 190 E7 007 40 000 4 66	B C B C
ATOM	9771 CG ASP	496	114. 938 58. 715 65. 154 1. 00 12. 87	B C
ATOM ATOM	9772 OD1 ASP 9773 OD2 ASP	496 496	113.849 59.297 65.330 1.00 12.34	B 0
ATOM	9774 C ASP			B 0
ATOM	9775 0 ASP	496	112.453 56.808 63.845 1.00 14.36	B C B 0
ATOM ATOM	9776 N ASN 9777 CA ASN	497 497	111. 820 58. 871 63. 234 1. 00 12. 87	B N
ATOM	9778 CB ASN	497	110. 460 58. 697 63. 717 1. 00 15. 91 109. 736 57. 666 62. 855 1. 00 16. 28	
ATOM	9779 CG ASN	497	109. 227 58. 255 61. 564 1. 00 20. 09	
ATOM ATOM	9780 OD1 ASN 9781 ND2 ASN	497 497	108.308 59.077 61.570 1.00 18.95	3 0
ATOM	9782 C ASN	497	109. 829 57. 853 60. 443 1. 00 19. 49 110. 373 58. 292 65. 193 1. 00 17. 71	
ATOM	9783 O ASN	497	109. 591 57. 420 65. 564 1. 00 19. 20 B	
ATOM ATOM	9784 N SER 9785 CA SER	498 498	111.179 58.924 66.035 1.00 18.90 B	N
ATOM	9786 CB SER	498	111. 147 58. 627 67. 458 1. 00 20. 75 112. 210 59. 454 68. 191 1. 00 20. 93	
ATOM	9787 OG SER	498	113. 491 58. 878 68. 037 1. 00 23. 33 B	
ATOM ATOM	9788 C SER 9789 O SER	498 498	109. 760 58. 956 68. 020 1. 00 20. 54 B	C
ATOM	9790 N ALA	499	109. 183 58. 184 68. 777 1. 00 20. 68 B 109. 238 60. 113 67. 637 1. 00 20. 46 B	
ATOM	9791 CA ALA	499	107. 935 60. 564 68. 087 1. 00 21. 87 B	N C
ATOM ATOM	9792 CB ALA 9793 C ALA	499 499	107. 577 61. 858 67. 391 1. 00 21. 73 B	C
ATOM	9794 O ALA	499 499	106. 859 59. 520 67. 822 1. 00 23. 85 B 106. 279 58. 961 68. 758 1. 00 25. 77 B	C
ATOM	9795 N LEU	500	106. 588 59. 262 66. 546 1. 00 23. 83 B	O N
ATOM ATOM	9796 CA LEU 9797 CB LEU	500 500	105. 568 58. 286 66. 176 1. 00 24. 31 B	C
ATOM	9798 CG LEU	500 500	105. 642 57. 958 64. 678 1. 00 22. 08 B 104. 618 56. 922 64. 201 1. 00 20. 35 B	C
ATOM	9799 CD1 LEU	500	104. 618 56. 922 64. 201 1. 00 20. 35 B 103. 200 57. 349 64. 570 1. 00 19. 30 B	C C
			· · · · · · · · · · · · · · · · · · ·	

	FIG. 4-201	(Continued)
ATOM 9811 N LYS ATOM 9812 CA LYS ATOM 9813 CB LYS ATOM 9814 CG LYS ATOM 9815 CD LYS ATOM 9816 CE LYS ATOM 9817 NZ LYS ATOM 9818 C LYS ATOM 9819 O LYS ATOM 9820 N MET ATOM 9821 CA MET ATOM 9821 CA MET ATOM 9822 CB MET ATOM 9823 CG MET ATOM 9824 SD MET ATOM 9825 CE MET ATOM 9826 C MET ATOM 9827 O MET ATOM 9827 O MET ATOM 9828 N LEU ATOM 9828 N LEU ATOM 9829 CA LEU ATOM 9830 CB LEU ATOM 9831 CG LEU ATOM 9831 CG LEU ATOM 9831 CG LEU ATOM 9832 CD1 LEU ATOM 9833 CD2 LEU ATOM 9834 C LEU ATOM 9835 O LEU ATOM 9836 N GLN ATOM 9836 N GLN ATOM 9837 CA GLN ATOM 9837 CA GLN ATOM 9838 CB GLN ATOM 9839 CG GLN ATOM 9839 CG GLN ATOM 9830 CB LEU ATOM 9831 CB CB CD ATOM 9831 CB CD ATOM 9834 C LEU ATOM 9834 C LEU ATOM 9835 O LEU ATOM 9836 N GLN ATOM 9837 CA GLN ATOM 9838 CB GLN ATOM 9839 CG GLN ATOM 9840 CD GLN ATOM 9841 OB1 GLN ATOM 9842 NE2 GLN ATOM 9844 O GLN ATOM 9845 N ASN ATOM 9845 N ASN ATOM 9846 CA ASN ATOM 9847 CB ASN	500 104.744 56.763 62.721 1.00 19.60 500 105.745 57.009 66.974 1.00 24.37 500 104.777 56.407 67.437 1.00 24.06 501 106.997 56.601 67.131 1.00 26.06 501 107.301 55.392 67.868 1.00 26.66 501 108.793 55.120 67.844 1.00 25.74 501 109.145 53.848 68.556 1.00 26.66 501 109.939 53.901 69.512 1.00 30.35 501 106.827 55.484 69.309 1.00 27.60 501 106.827 55.484 69.309 1.00 27.99 502 107.011 56.645 69.924 1.00 28.69 502 106.591 56.819 71.301 1.00 33.97 502 106.507 58.484 73.239 1.00 35.56 502 106.991 59.822 73.766 1.00 36.56 502 106.997 59.	C C O N C C C O O C O N C C C C C N C O N C C C S C C O N C C C C C O N C C C C O N C C C C
ATOM 9833 CD2 LEU 5 ATOM 9834 C LEU 5 ATOM 9835 O LEU 5 ATOM 9836 N GLN 5 ATOM 9837 CA GLN 5 ATOM 9838 CB GLN 5 ATOM 9839 CG GLN 5 ATOM 9840 CD GLN 5 ATOM 9841 OB1 GLN 5 ATOM 9842 NE2 GLN 5 ATOM 9842 NE2 GLN 5 ATOM 9843 C GLN 5 ATOM 9844 O GLN 5 ATOM 9845 N ASN 5 ATOM 9846 CA ASN 50	04 101.696 54.750 66.106 1.00 28.10 04 103.508 53.300 65.143 1.00 27.73 04 102.581 52.998 69.986 1.00 29.56 04 101.880 51.991 70.016 1.00 27.71 05 103.458 53.291 70.938 1.00 31.52 05 103.641 52.425 72.096 1.00 33.96 05 104.829 52.915 72.927 1.00 36.96 05 106.167 52.836 72.200 1.00 42.44 05 106.652 51.408 71.996 1.00 43.93 05 106.581 50.925 70.758 1.00 45.80 05 106.581 50.925 70.758 1.00 45.49 05 102.375 52.393 72.960 1.00 33.38 05 102.104 51.400 73.634 1.00 32.77 06 10.607 53.482 72.928 1.00 32.38 <	B C B O B N B C B C B C B C B C B C B C B C B C B C

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					FIC	G. 4-	202			(Continued)
ATOM	9849		ASN	506	101.426	55.608	75. 799	1.00 41.09	В	0
ATOM		ND2		506	101.703	56. 796	73.903	1.00 39.00	В	N
ATOM	9851	C	ASN	506	99. 208	52.933	72.936	1.00 30.32	В	C
ATOM	9852	0	ASN	506	98.058	52. 995	73. 377	1.00 30.93	В	0
ATOM	9853	N	VAL	507	99. 516	52.305	71.803	1.00 26.94	В	N
ATOM	9854	CA	VAL	507	98. 497	51.664	70.974	1.00 25.15	В	C
ATOM	9855	CB	VAL		98. 456	52. 293	69.545	1.00 23.88	В	C
ATOM	9856		VAL	507	97. 287	51.730	68.755	1.00 21.31	В	C .
ATOM	9857		VAL	507	98. 344	53. 811	69.633	1.00 22.11	В	C
ATOM ATOM	9858	C	VAL	507	98.717	50. 164	70.825	1.00 25.62	В	C
ATOM	9859 9860	O N	VAL GLN	507 508	99. 838 97. 639	49.676	70. 945 70. 567	1.00 26.78	B B	0
ATOM	9861	CA	GLN	508	97. 730	49. 432 47. 992	70. 381	1.00 25.89 1.00 25.14	В	N
ATOM	9862	CB	GLN	508	96. 486	47. 281	70. 351	1.00 23.14	В	C C
ATOM	9863	CG	GLN	508	96. 322	47. 397	72. 422	1.00 21.32	В	C
ATOM	9864	CD	GLN	508	95. 190	46. 543	72. 958	1.00 30.81	В	Č .
ATOM	9865	0E1		508	95. 208	45. 312	72.836	1.00 31.32	B	Ö
ATOM	9866		GLN	508	94. 199	47. 190	73. 561	1.00 29.92	B	Ň
ATOM	9867	C	GLN	508	97.869	47. 740	68.899	1.00 23.65	B	Ċ
ATOM	9868	0	GLN	508	96.944	47.277	68. 241		B	ŏ
ATOM	9869	N	MET	509	99.046	48.063	68.385	1.00 23.78	B	Ň
ATOM	9870	CA	MET	509	99.347	47.895	66.980	1.00 23.48	В	Č
ATOM	9871	CB	MET	509	100.667	48.578	66.655	1.00 23.41	В	C
ATOM	9872	CG	MET	509	100.586	50.070	66.782	1.00 26.19	В	С
ATOM	9873	SD	MET	509	99.279	50.681	65.719	1.00 28.03	В	S
ATOM	9874	CE	MET	509	100. 207	50.994	64. 209	1.00 25.78	В	C
ATOM	9875	C	MET	509	99.425	46.440	66. 579	1.00 23.44	В	C
ATOM	9876	0	MET	509	99.902	45. 599	67. 343	1.00 24.15	В	0
ATOM	9877	N	PRO	510	98. 951	46. 121	65.365	1.00 22.69	В	N
ATOM	9878	CD	PRO	510	98.308	47.027	64. 395	1.00 22.87	В	C
ATOM	9879	CA	PRO	510	98. 974	44. 751	64.854	1.00 21.97	В	. C
ATOM ATOM	9880	CB CG	PRO	510 510	97. 987	44. 807	63.701	1.00 22.62	В	C
ATOM	9881 9882	C	PRO	510	98. 248	46. 171	63.141	1.00 22.72	В	C
ATOM	9883	0	PRO PRO	510 510	100. 381 101. 249	44. 434		1.00 21.20	. В	C
ATOM	9884	N	SER	511	101. 249	45. 301 43. 188	64. 353 63. 997	1.00 19.97 1.00 22.07	В	0 N
ATOM	9885	CA	SER	511	100.003	43. 188	63. 521	1.00 22.07	B B	N
ATOM	9886	CB	SER	511	102. 481	41.654	64. 392	1.00 23.02	В	C C
ATOM	9887	OG	SER	511	101.653	40.500	64. 358	1.00 25.03	В	0
ATOM	9888	Č	SER	511	101.773	42. 299	62. 094	1.00 23.35	В	C
ATOM	9889	Ö	SER	511	100.659	42. 168	61.583	1.00 24.92	В	0
ATOM	9890	Ň	LYS	512	102.906	42.035	61.458	1.00 22.83	В	N
ATOM	9891		LYS	512	102.916	41.556	60.094	1.00 22.46	B	C
ATOM	9892	CB	LYS	512	103.490	42.615	59.168	1.00 21.81	B	č
ATOM	9893	CG	LYS	512	103. 494	42. 209	57. 705	1.00 23.24	B	č
ATOM	9894	CD	LYS	512	103.820	43. 411	56.851	1.00 24.28	B	č
ATOM	9895	CE	LYS	512	103.824	43.080	55. 393	1.00 23.13	B	č
ATOM	9896		LYS	512	104.160	44. 299	54.622	1.00 24.52	В	N
ATOM	9897	С	LYS	512	103. 742	40. 289	59.993	1.00 22.87	В	C.

					FIG. 4-203	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9898 9899 9900 9901 9902 9903	N CA CB CG CD	LYS LYS LYS LYS LYS LYS	512 513 513 513 513	104. 803 40. 180 60. 585 1. 00 23. 26 B 103. 235 39. 331 59. 235 1. 00 24. 10 B 103. 910 38. 069 59. 039 1. 00 24. 49 B 103. 046 36. 923 59. 566 1. 00 25. 52 B 103. 522 35. 537 59. 148 1. 00 26. 69 B 102. 493 34. 471 59. 522 1. 00 30. 85 B	O N C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9904 9905 9906 9907 9908 9909 9910	CE NZ C O N CA CB CG	LYS LYS LYS LYS LEU LEU LEU LEU	513 513 513 513 514 514 514	102. 805 33. 124 58. 866 1. 00 33. 37 B 104. 131 32. 573 59. 287 1. 00 36. 04 B 104. 143 37. 888 57. 552 1. 00 25. 44 B 103. 196 37. 871 56. 763 1. 00 27. 00 B 105. 409 37. 771 57. 171 1. 00 24. 62 B 105. 775 37. 561 55. 783 1. 00 22. 99 B 106. 870 38. 536 55. 380 1. 00 22. 15 B 107. 307 38. 465 53. 925 1. 00 21. 19 B	C N C O N C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9912 9913 9914 9915 9916 9917 9918 9919		LEU LEU LEU ASP ASP ASP	514 514 514 514 515 515	106. 125 38. 790 53. 029 1.00 19. 85 B 108. 438 39. 435 53. 701 1.00 18. 42 B 106. 292 36. 132 55. 708 1.00 24. 30 B 107. 123 35. 725 56. 519 1.00 24. 87 B 105. 804 35. 361 54. 747 1.00 25. 31 B 106. 233 33. 975 54. 634 1.00 26. 30 B 105. 599 33. 156 55. 757 1.00 28. 58 B	C C O N C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9920 9921 9922 9923 9924 9925 9926	0D1	ASP ASP ASP ASP ASP PHE PHE	515 515 515 515 515 516 516 516	106. 403 31. 929 56. 108 1. 00 30. 08 B 107. 209 31. 474 55. 272 1. 00 31. 89 B 106. 216 31. 409 57. 224 1. 00 33. 36 B 105. 805 33. 414 53. 282 1. 00 26. 17 B 105. 343 34. 157 52. 417 1. 00 26. 57 B 105. 940 32. 104 53. 103 1. 00 25. 46 B 105. 571 31. 496 51. 838 1. 00 25. 82 B 106. 792 31. 384 50. 930 1. 00 23. 83 B	C 0 0 C 0 N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9927 9928 9929 9930 9931 9932 9933	CG CD1 CD2 CE1 CE2 CZ CZ	PHE PHE PHE PHE PHE PHE PHE	516 516 516 516 516 516 516	106. 792 31. 384 50. 930 1. 00 23. 83 B 107. 811 30. 395 51. 413 1. 00 22. 29 B 108. 896 30. 808 52. 176 1. 00 22. 68 B 107. 678 29. 042 51. 119 1. 00 21. 58 B 109. 836 29. 885 52. 642 1. 00 21. 89 B 108. 609 28. 113 51. 579 1. 00 21. 19 B 109. 689 28. 536 52. 342 1. 00 20. 70 B 104. 955 30. 117 51. 954 1. 00 26. 95 B	C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	9934 9935 9936 9937 9938 9939	CG1 CD1	ILE	516 517 517 517 517 517	105. 063 29. 452 52. 980 1. 00 28. 94 B 104. 307 29. 707 50. 872 1. 00 27. 35 B 103. 697 28. 398 50. 755 1. 00 28. 12 B 102. 155 28. 470 50. 729 1. 00 26. 53 B 101. 645 29. 073 52. 016 1. 00 27. 39 B 101. 682 29. 296 49. 537 1. 00 27. 43 B 100. 175 29. 486 49. 486 1. 00 26. 37 B	O N C C C C
ATOM ATOM ATOM ATOM ATOM ATOM	9941 9942 9943 9944 9945 9946	C O N CA CB CG2	ILE ILE ILE ILE ILE ILE	517 517 518 518 518 518	104. 202 27. 896 49. 411 1.00 30.13 B 104. 575 28. 697 48. 551 1.00 29. 21 B 104. 239 26. 581 49. 228 1.00 33. 16 B 104. 709 26. 029 47. 969 1.00 36. 01 B 105. 680 24. 867 48. 190 1.00 36. 84 B 106. 133 24. 311 46. 845 1.00 36. 94 B	C O N C C C

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					FIC	4 - 204	•		(Continued)
ATOM	9947		ILE	518		349 49.000	1.00 38.21	В	C
ATOM	9948	CD1		518		296 49.169	1.00 40.77	В	C
ATOM ATOM	9949 9950	C 0	ILE ILE	518		534 47.114	1.00 37.38	В	C
ATOM	9951	N	LEU	518 519		000 47.624 730 45.808	1.00 38.97	В	0
ATOM	9952	CA	LEU	519		294 44.863	1.00 39.11 1.00 40.68	B B	N C
ATOM	9953	CB	LEU	519		461 44.474	1.00 40.08	В	C C
ATOM	9954	CG	LEU	519		144 45.612	1.00 39.82	В	C
ATOM	9955		LEU	519		205 45.045	1.00 39.14	В	C
ATOM	9956		LEU	519		107 46.381	1.00 40.51	B	č
ATOM	9957	C	LEU	519		763 43.637	1.00 42.22	B	č
ATOM	9958	0	LEU	519		524 42.910	1.00 42.60	B	ŏ
ATOM	9959	N	ASN	520		453 43.419	1.00 43.53	В	N
ATOM	9960	CA	ASN	520		824 42.285	1.00 44.57	В	C
ATOM	9961	CB	ASN	520		337 40.964	1.00 46.39	В	C
ATOM	9962	CG	ASN	520		726 40.639	1.00 48.97	В	C
ATOM	9963		ASN	520		634 41.498	1.00 50.54	В	0 .
ATOM	9964		ASN	520		312 39.386	1.00 50.46	В	N
ATOM ATOM	9965	C	ASN	520		114 42.316	1.00 44.13	В	C
ATOM	9966 9967	O N	ASN GLU	520 521		637 41.348	1.00 44.64	В	0
ATOM	9968	CA	GLU	521	106. 097 22. 107. 536 23.	791 43.431 012 43.562	1.00 44.10	В	N
ATOM	9969	CB	GLU	521		387 42.368	1.00 45.15 1.00 49.07	В	C
ATOM	9970	CG	GLU	521		642 42.339	1.00 49.07	B B	C
ATOM	9971	CD	GLU	521		274 41.004	1.00 58.04	В	C
ATOM	9972		GLU	521	110. 307 21.		1.00 59.07	В	Õ
ATOM	9973		GLU	521		176 40.361	1.00 59.78	B	ő
ATOM	9974	C	GLU	521	107. 922 24.		1.00 42.18	B	č
ATOM	9975	0	GLU	521		810 44.072	1.00 42.85	B	. 0
ATOM	9976	Ŋ	THR	522	107.014 25.		1.00 38.59	В	N
ATOM	9977	CA	THR	522	107. 314 26.		1.00 34.63	В	C .
ATOM	9978	CB	THR	522	106.605 27.		1.00 34.21	В	C
ATOM	9979		THR	522	107. 109 27.		1.00 34.20	В	0
ATOM	9980		THR	522	106. 866 29.		1.00 33.69	В	C
ATOM ATOM	9981 9982	C	THR	522	106.959 27.4		1.00 32.83	В	C
ATOM	9983	O N	THR LYS	522 523	106.028 27.		1.00 32.75	В	0
ATOM	9984	CA	LYS	523	107. 727 28. 4 107. 559 29. 3		1.00 31.06	В	N
ATOM	9985	CB	LYS	523	107. 559 29. 1 108. 940 29. 4		1.00 29.30	В	C
ATOM	9986	CG	LYS	523	108. 934 30.		1.00 29.00 1.00 31.42	B B	C
ATOM	9987	CD	LYS	523	110.344 30.		1.00 31.42	В	C C
ATOM	9988	CE	LYS	523	111.045 29.5		1.00 32.07	В	C
ATOM	9989	NZ	LYS	523	112.388 29.		1.00 35.72	В	N
ATOM	9990	C	LYS	523	106.819 30.		1.00 28.56	B	Č
ATOM	9991	0	LYS	523	107. 256 31.		1.00 29.36	B	ŏ
ATOM	9992	N	PHE	524	105.692 30.7		1.00 25.40	В	Ň
ATOM	9993	CA	PHE	524	104.912 31.9		1.00 22.61	В	C
ATOM	9994	CB	PHE	524	103.529 31.6		1.00 22.69	В	C
ATOM	9995	CG	PHE	524	103.565 31.1	136 44.516	1.00 21.75	В	C

		FIG. 4-205	(Continued)
ATOM 9997 CD2 ATOM 9998 CE1 ATOM 9999 CE2 ATOM 10000 CZ ATOM 10001 C ATOM 10002 O ATOM 10003 N ATOM 10004 CA ATOM 10005 CB ATOM 10006 CG ATOM 10007 CD2 ATOM 10008 CE2 ATOM 10009 CE3 ATOM 10010 CD1 ATOM 10011 NE1 ATOM 10012 CZ2 ATOM 10013 CZ3 ATOM 10014 CH2 ATOM 10015 C ATOM 10015 C ATOM 10016 O ATOM 10017 N ATOM 10018 CA ATOM 10019 CB ATOM 10010 CD1 ATOM 10019 CB ATOM 10020 CG ATOM 10021 CD1 ATOM 10020 CG ATOM 10020 CG ATOM 10021 CD1 ATOM 10022 CE1 ATOM 10022 CE1 ATOM 10023 CD2 ATOM 10024 CE2 ATOM 10025 CZ ATOM 10025 CZ ATOM 10026 OH ATOM 10027 C ATOM 10027 C ATOM 10028 O ATOM 10029 N ATOM 10029 N ATOM 10030 CA ATOM 10030 CA ATOM 10031 CB ATOM 10030 CA ATOM 10040 CB ATOM 10040 CB ATOM 10042 SD ATOM 10043 CE	TRP 525 TRP 526 TYR 527 SLN 528 SET 528 SET 528 SET 528	103. 626 29. 773 44. 247 1. 00 22. 19 103. 541 32. 031 43. 448 1. 00 22. 40 B 103. 662 29. 306 42. 935 1. 00 22. 45 B 103. 576 31. 579 42. 131 1. 00 22. 70 B 103. 637 30. 213 41. 871 1. 00 22. 70 B 104. 765 32. 593 47. 890 1. 00 20. 73 B 104. 416 31. 941 48. 875 1. 00 19. 19 B 105. 016 33. 892 47. 950 1. 00 18. 35 B 104. 950 34. 600 49. 216 1. 00 17. 31 B 106. 059 35. 646 49. 274 1. 00 16. 81 B 107. 442 35. 092 49. 191 1. 00 16. 14 B 108. 393 35. 031 50. 253 1. 00 14. 70 B 108. 366 35. 411 51. 602 1. 00 14. 22 B 108. 062 34. 560 48. 086 1. 00 15. 59 B 109. 344 34. 176 48. 403 1. 00 14. 99 B 110. 722 34. 247 50. 508 1. 00 17. 11 B 109. 506 35. 204 52. 381 1. 00 14. 40 B 110. 668 34. 627 51. 829 1. 00 15. 16 B 103. 630 35. 280 49. 554 1. 00 17. 78 B 102. 880 35. 719 48. 675 1. 00 17. 96 B 103. 361 35. 368 50. 849 1. 00 15. 16 B 101. 053 35. 030 51. 652 1. 00 19. 10 B 101. 369 34. 076 52. 778 1. 00 21. 69 B 101. 132 34. 422 54. 110 1. 00 23. 52 B 101. 416 33. 531 55. 146 1. 00 22. 69 B 101. 902 32. 817 52. 509 1. 00 24. 93 B 102. 189 31. 922 53. 527 1. 00 24. 93 B 102. 189 31. 922 53. 527 1. 00 24. 93 B 102. 189 31. 922 53. 527 1. 00 24. 93 B 102. 189 31. 922 53. 527 1. 00 24. 93 B 102. 189 31. 922 53. 527 1. 00 24. 93 B 101. 945 32. 280 54. 840 1. 00 17. 78 B 102. 847 34. 849 54. 840 1. 00 28. 49 B 102. 235 31. 370 55. 830 1. 00 31. 26 B 101. 901 38. 417 54. 250 1. 00 17. 09 B 101. 903 44. 347 54. 250 1. 00 17. 09 B 101. 903 38. 417 54. 250 1. 00 17. 09 B 101. 904 32. 280 54. 840 1. 00 28. 49 B 102. 235 31. 370 55. 830 1. 00 31. 26 B 101. 945 32. 280 54. 840 1. 00 28. 49 B 102. 189 31. 992 52. 509 1. 00 16. 76 B 103. 600 36. 536 53. 877 1. 00 26. 65 B 101. 945 32. 280 54. 840 1. 00 23. 52 B 101. 462 40. 740 53. 234 1. 00 15. 60 B 101. 901 38. 417 54. 250 1. 00 17. 09 B 101. 901 38. 417 54. 250 1. 00 17. 09 B 101. 901 38. 417 54. 250 1. 00 17. 09 B 101. 901 38. 417 54. 250 1. 00 17. 09 B 100. 576 38. 630 57. 776 1. 00 23. 43 B 99. 519 38. 439 54. 269 1. 00 20. 94 B SUBSTITUTE SHEET (RULE 26)	CCCCCONCCCCCCNCCCCONCCCCCCCCCCONCCCCCONCCONCCCCSCC

					FIG. 4	206		(Continued)
ATOM	10045	0	MET	528		•	.	0
ATOM	10045	N	ILE	529	100. 471 39. 845 98. 432 40. 554	58. 720 1. 00 21. 07 58. 100 1. 00 19. 01	B B	0 N
ATOM	10047	CA	ILE	529	98. 428 41. 614	59. 082 1. 00 18. 89	В	N C
ATOM	10048	CB	ILE	529	97.718 42.860	58.540 1.00 16.80	В	Č
ATOM	10049		2 ILE	529	97.656 43.937	59. 615 1. 00 13. 98	В	č
ATOM	10050		ILE	529	98.469 43.368	57. 296 1. 00 15. 06	B	Č
ATOM	10051		ILE	529	99. 934 43. 701	57. 537 1. 00 11. 03	В	C
ATOM	10052	C	ILE	529	97. 656 40. 969	60. 225 1. 00 20. 15	В	C
ATOM	10053	0	ILE	529	96. 457 40. 720	60. 124 1. 00 20. 94	В	0
ATOM	10054	N	LEU	530	98. 359 40. 653	61. 302 1. 00 21. 19	В	N
ATOM ATOM	10055 10056	CA CB	LEU LEU	530 530	97. 717 39. 985	62. 420 1. 00 21. 61	В	C
ATOM	10057	CG	LEU	530	98. 649 38. 907 99. 086 37. 875	62. 976 1. 00 19. 85 61. 931 1. 00 19. 34	В	C
ATOM	10058		LEU	530	100. 238 37. 027	61. 931 1. 00 19. 34 62. 461 1. 00 20. 33	B B	C
ATOM	10059		LEU	530	97. 897 37. 010	61.562 1.00 19.04	В	C C
ATOM	10060	C	LEU	530	97. 294 40. 930	63. 521 1. 00 22. 34	В	Č .
ATOM	10061	0	LEU	530	98.006 41.878	63. 854 1. 00 23. 45	В	Õ
ATOM	10062	N	PR0	531	96.104 40.697	64.088 1.00 23.19	В	Ň
ATOM	10063	CD	PR0	531	95. 105 39. 684	63. 711 1. 00 22. 71	B	Ċ
ATOM	10064	CA	PRO	531	95.600 41.545	65.169 1.00 24.33	В	C
ATOM	10065	CB	PRO	531	94.188 41.002	65.404 1.00 22.74	В	C
ATOM	10066	CG	PRO	531	94. 276 39. 588	64.967 1.00 23.03	В	C
ATOM ATOM	10067 10068	C	PRO	531	96. 490 41. 438	66. 407 1. 00 25. 18	В	C
ATOM	10068	O N	PRO PRO	531 532	97. 244 40. 478	66. 562 1. 00 24. 64	В	0
ATOM	10003	CD	PRO	532	96. 424 42. 433 95. 502 43. 581	67. 300 1. 00 26. 64	В	N
ATOM	10071	CA	PRO	532	95. 502 43. 581 97. 246 42. 397	67. 326 1. 00 25. 36 68. 513 1. 00 27. 91	В	C
ATOM	10072	CB	PRO	532	96. 868 43. 698	68.513 1.00 27.91 69.216 1.00 27.08	В	C
ATOM	10073	CG	PRO	532	95. 443 43. 897	68. 793 1. 00 26. 25	В В	C C C C
ATOM	10074	C	PRO	532	96. 945 41. 160	69. 369 1. 00 29. 25	В	C
ATOM	10075	0	PR0	532	95.865 40.579	69. 279 1. 00 29. 62	В	Ŏ
ATOM	10076	N	HIS	533	97. 909 40. 756	70.187 1.00 30.65	B	Ň
ATOM	10077	CA	HIS	. 533	97. 738 39. 602	71.061 1.00 31.99	B	Ĉ.
ATOM	10078	CB	HIS	533	96. 749 39. 945	72.172 1.00 32.50	В	Č
ATOM	10079	CG	HIS	533	96. 981 41. 293	72. 783 1. 00 35. 12	В	C
ATOM	10080		HIS	533	96. 168 42. 370	72. 903 1. 00 36. 18	В	C
ATOM ATOM	10081		HIS	533	98. 181 41. 653	73. 358 1. 00 35. 49	В	N
ATOM	10082 10083		HIS HIS	533 533	98.096 42.892	73. 807 1. 00 36. 37	В	C
ATOM	10084	C	HIS	533	96. 885 43. 350 97. 249 38. 382	73. 544 1. 00 37. 01 70. 286 1. 00 33. 21	В	N
ATOM	10085	ŏ	HIS	533	97. 249 38. 382 96. 447 37. 590		В	C
ATOM	10086	Ň	PHE	534	97. 739 38. 243	70. 791 1. 00 32. 78 69. 058 1. 00 33. 50	B B	O N
ATOM	10087	CA	PHE	534	97. 374 37. 125	68. 200 1. 00 34. 63	В	C
ATOM	10088	CB	PHE	534	98. 283 37. 085	66.970 1.00 32.35	В	Č
ATOM	10089	CG	PHE	534	97.997 35.942	66.041 1.00 32.06	В	Č
ATOM	10090		PHE	534	96. 790 35. 871	65. 354 1. 00 32. 10	B	č
ATOM	10091		PHE	534	98. 936 34. 938	65. 848 1. 00 32. 66	В	C
ATOM	10092	CE1		534	96. 522 34. 819	64.486 1.00 31.59	В	C
ATOM	10093	CE2	rne	534	98. 679 33. 879	64. 982 1. 00 32. 91	В	C

					FIG. 4-207	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10094 10095 10096 10097 10098 10099 10100 10101 10105 10106 10107 10108 10110 10111 10111 10111 10111 10111 10115 10116 10117 10118 10119 10120 10121 10121	C O N CA CB CG OD OD C C O N CA CB CG CD N CA CB CG CD N CA CB CC	PHE PHE PHE ASP ASP ASP ASP LYS	534 534 535 535 535 535 535 536 536 536 536 536	97. 469	C C C O N C C C C C N C O N C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C C O N C O N C C O N C C O N C C O N C O N C O N C O N C O N C C O N C O
ATOM ATOM ATOM ATOM ATOM	10122 10123 10124 10125 10126 10127 10128 10130 10131 10132 10133 10134 10135 10136 10137 10138 10139 10140 10141	CB CG CD CE NZ C O N CA CB CC CD CE NZ C C O N CA CB CC CD CE NZ C C O N CA CB CC CD CC CD C CD C CD C CD C CD C	LYS		91. 629	C C C C C C C C C C C C C C C C C C C

						•				
					FIC	G. 4-	208			(Continued)
ATOM	10143	CE1	TYR	540	93. 321	34. 708	66. 130	1.00 21.65	В	С
ATOM	10144		TYR	540	91.810	36. 863	65. 257	1.00 22.89	В	č
ATOM	10145		TYR	540	92. 507	36. 955	66. 449	1.00 22.77	В	Č
ATOM	10146	CZ	TYR	540	93. 261	35. 875	66. 881	1.00 22.87	В	C C C
ATOM	10147	OH	TYR	540	93. 950	35. 965	68.062	1.00 23.97	В	ŏ
ATOM	10148	C	TYR	540	89. 335	34. 694	61.749	1.00 23.62	В	Č
ATOM	10149	Ö	TYR	540	89. 670	33. 842	60. 925	1.00 23.93	В	ŏ
ATOM	10150	N	PRO	541	88. 457	35.660	61.452	1.00 21.89	В	N N
ATOM	10151	CD	PRO	541	87. 820	36. 667	62.320	1.00 21.33	В	C
ATOM	10152	CA	PRO	541	87. 917	35. 719	60.095	1.00 20.52	В	Č
ATOM	10152	CB	PRO	541	86.770	36. 717	60. 228	1.00 20.32	В	C
ATOM	10153	CG	PRO	541	87. 243	37. 629	61.317	1.00 20.36	В	Č
ATOM	10155	C	PRO	541	89. 077	36. 266	59.276	1.00 20.30	В	C
ATOM	10156	Õ	PRO	541	90. 026	36. 799	59.841	1.00 19.00	В	0
ATOM	10157	Ň	LEU	542	89.028	36. 147	57.961	1.00 19.38	В	N
ATOM	10158	CA	LEU	542	90.133	36.655	57.169	1.00 13.33	В	C
ATOM	10159	CB	LEU	542	91.027	35. 483	56. 741	1.00 18.21	В	Č
ATOM	10160	ČĞ	LEU	542	92. 215	35. 768	55.816	1.00 10.30	В	Č
ATOM	10161		LEU	542	93. 296	34. 721	56.025	1.00 17.89	В	č
ATOM	10162		LEU	542	91.741	35. 775	54.374	1.00 19.31	В	č
ATOM	10163	C	LEU	542	89. 677	37. 458	55.954	1.00 17.31	В	Č
ATOM	10164	ő	LEU	542	88. 720	37. 087	55. 282	1.00 18.08	В	Ö
ATOM	10165	Ň	LEU	543	90.368	38. 564	55.694	1.00 14.81	В	N N
ATOM	10166	CA	LEU	543	90.075	39.430	54.559	1.00 13.79	В	Č
ATOM	10167	CB	LEU	543	89.816	40.872	55.015	1.00 12.33	В	č
ATOM	10168	CG	LEU	543	89. 568	41.892	53.886	1.00 13.71	. В	č
ATOM	10169		LEU	543	88. 317	41.497	53.113	1.00 9.91	В	Č
ATOM	10170		LEU	543	89. 409	43. 294	54.454	1.00 11.87	В	č
ATOM	10171	C	LEU	543	91. 273	39. 415	53.620	1.00 14.35	В	Č
ATOM	10172	Ŏ	LEU	543	92.349	39. 893	53.966	1.00 14.00	В	Ŏ
ATOM	10173	Ň	LEU	544	91.091	38. 866	52.428	1.00 15.02	B	Ň
ATOM	10174	CA	LEU	544	92. 191	38. 807	51.480	1.00 16.19	В	Č .
ATOM	10175	CB	LEU	544	92.006	37. 609	50.539	1.00 16.34	В	Č
ATOM	10176	CG	LEU	544	93. 163	37. 231	49.608	1.00 14.93	В	č
ATOM	10177		LEU	544	94. 345	36. 752	50.429	1.00 15.36	В	č
ATOM	10178		LEU	544	92.713	36.128	48.654	1.00 15.79	В	Č.
ATOM	10179	C	LEU	544	92. 276	40. 109	50.679	1.00 16.49	B	Č
ATOM	10180	0	LEU	544	91.437	40.374	49.819	1.00 17.02	В	ŏ
ATOM	10181	Ň	ASP	545	93. 280	40. 925	50.997	1.00 15.13	B	Ň
ATOM	10182	CA	ASP	545	93. 515	42.186	50.306	1.00 14.91	В	
ATOM	10183	CB	ASP	545	94.479	43.069	51.117	1.00 15.71	B	C C
ATOM	10184	CG	ASP	545	94. 703	44. 434	50. 483	1.00 15.88	B	č
ATOM	10185		ASP	545	94. 285	44.641	49.324	1.00 14.36	B	Ŏ .
ATOM	10186	OD2		545	95. 304	45.304	51.144	1.00 15.41	B	Ŏ
ATOM	10187	C	ASP	545	94. 175	41.757	49.004	1.00 14.61	B	č
ATOM	10188	0	ASP	545	95. 235	41.135	49.014	1.00 13.17	B	ŏ
ATOM	10189	N	VAL	546	93. 567	42.098	47. 881	1.00 15.03	B	Ň
ATOM	10190	CA	VAL	546	94. 116	41.667	46.614	1.00 17.39	B	Č
ATOM	10191	CB	VAL	546	93. 199	40.579	46.014	1.00 19.44	B	č

					FIC	. 1	209			(Con	itinued)
					FIG	'• '1	_2 0 9				
ATOM	10192	CG:	1 VAL	546	93.717	40.124	44.647	1.00 17.8	7 B	С	
ATOM	10193	CG	2 VAL	546	93.109	39.410		1.00 20.9		č	
ATOM	10194	C	VAL	546	94.343	42.722		1.00 17.0		č	
ATOM	10195	0	VAL	546	93.601	43.694		1.00 18.1		ŏ	
ATOM	10196	N	TYR	547	95. 391	42.519		1.00 15.7		Ň	
ATOM	10197		TYR	547		43.378		1.00 14.9		Ĉ	
ATOM	10198	CB	TYR	547	96.838	44.335		1.00 12.5		Č	
ATOM	10199		TYR	547	97.008	45. 241	42.622	1.00 12.8		Ç.	
ATOM	10200		I TYR	547		45.063	41.727	1.00 12.0		Č	
ATOM	10201		TYR	547	98. 165	45.839	40.578	1.00 9.9		Č	
ATOM	10202		? TYR	547		46.226	42.331	1.00 11.83		C	
ATOM	10203		? TYR	547		47.002	41.183	1.00 8.65		C	
ATOM	10204	CZ	TYR	547		46.804	40.314	1.00 10.60		C	
ATOM	10205	OH	TYR	547		47. 573	39.179	1.00 12.10) B	0	
ATOM	10206	C	TYR	547			42.485	1.00 13.60) B	C	
ATOM	10207	0	TYR	547		42. 205	41.548	1.00 13.39		0	
ATOM	10208	N	ALA	548		41.763	42.608	1.00 13.66		N	
ATOM	10209	CA	ALA	548		40. 730	41.672	1.00 14.14		C	
ATOM	10210	CB	ALA	548		39. 518	41.807	1.00 11.57		C	
ATOM	10211	C	ALA	548		41.105	40. 207	1.00 13.67		C	
ATOM	10212	0	ALA	548		40. 234	39. 340	1.00 14.21		0	
ATOM ATOM	10213	N	GLY	549		42. 386	39. 913	1.00 13.87		N	
ATOM	10214 10215	CA	GLY	549		42. 765	38. 524	1.00 12.26		C	
ATOM	10215	C 0	GLY GLY	549		42. 209	38. 046	1.00 12.16		C	
ATOM	10210	N	PRO	549		41.717	38. 855	1.00 12.33		0	
ATOM	10211	CD	PRO	550 550		42. 256	36. 739	1.00 13.98		N	
ATOM	10219	CA	PRO	550 550		42.760	35.644	1.00 12.99		C	
ATOM	10213	CB	PRO	550 550		41. 736	36. 217	1.00 13.32		C	
ATOM	10221	CG	PRO	550		42.007 42.015	34. 721	1.00 14.56	В	C	
ATOM	10222	C	PRO	550		42.459	34. 473 36. 832	1.00 14.10		C	
ATOM	10223	ŏ	PRO	550		43. 683	36. 785	1.00 13.86 1.00 13.45		C	
ATOM	10224	Ň	CYS	551		41.694	30. 185 37. 405	1.00 13.45		0	
ATOM	10225	ĊA	CYS	551		42. 244	38. 027	1.00 14.79	B B	N C	
ATOM	10226	CB	CYS	551		43. 139	37. 036	1.00 13.31		_	
ATOM	10227	SG	CYS	551		43. 567	37. 543	1.00 17.09		C S	
ATOM	10228	C	CYS	551		43.018	39. 312	1.00 16.05	В	S C	•
ATOM	10229	0	CYS	551		43. 938	39. 702	1.00 15.36	В	0	
ATOM	10230	N	SER	552		12.631	39. 976	1.00 15.15	В	N	
ATOM	10231	CA	SER	552		13. 268	41. 229	1.00 14.65	В	C	
ATOM	10232	CB	SER	552		13. 149	41. 425	1.00 14.00	В	Č	•
ATOM	10233	0G	SER	552		11.789		1.00 14.39	В	0	
ATOM	10234	C	SER	552		12.608		1.00 15.21	В	C	
ATOM	10235	0	SER	552		11.585		1.00 15.34	В	Õ	
ATOM	10236	N	GLN	553		3. 201		1.00 14.73	B	N	
ATOM	10237	CA	GLN	553		2.647		1.00 14.31	B	Ċ	
ATOM	10238	CB	GLN	553		3.017		1.00 13.21	B	č	
ATOM	10239	CG	GLN	553		2. 332		1.00 15.05	B	Č	
ATOM	10240	CD	GLN	553		2. 585		1.00 15.66	В	Č	
					elipetitiite (CHEET					

						٠	٠.			(Cor	ntinued)
* •					FIC	G. 4`-	2 1 0			(001	ionnaca)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10241 10242 10243 10244 10245 10246 10247		GLN GLN GLN LYS LYS LYS	553	107. 812 108. 138 102. 921 103. 148 102. 031 101. 284 99. 817	43. 686 41. 556 43. 166 44. 295 42. 344 42. 734 42. 318	46. 400 45. 773 46. 012 46. 434 46. 568 47. 754 47. 633	1.00 16.56 1.00 15.50 1.00 14.58 1.00 14.77 1.00 14.78 1.00 16.57 1.00 17.81	B B B B B	0 N C 0 N C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10248 10249 10250 10251 10252 10253 10254 10255	CG CD CE NZ C O N CA	LYS LYS LYS LYS LYS ALA ALA	554 554 554 554 554 554 555	99. 031 99. 047 98. 228 96. 769 101. 890 101. 424 102. 939 103. 622	43. 142 44. 612 44. 902 44. 771 42. 148 42. 429 41. 350 40. 730	46. 630 47. 000 48. 261 48. 035 49. 024 50. 124 48. 866 50. 004	1.00 18.63 1.00 18.55 1.00 18.33 1.00 13.33 1.00 16.05 1.00 17.37 1.00 15.91 1.00 15.84	B B B B B B	C C C N C O N C	
ATOM ATOM ATOM ATOM ATOM ATOM	10256 10257 10258 10259 10260 10261 10262	CB C O N CA CB	ALA ALA ASP ASP ASP ASP	555 555 555 556 556 556	103. 656 105. 041 105. 954 105. 233 106. 571 106. 801 105. 750	39. 210 41. 246 40. 691 42. 304 42. 854 44. 085 45. 159	49. 833 50. 142 49. 539 50. 924 51. 134 50. 243 50. 430	1.00 15.51 1.00 14.91 1.00 15.57 1.00 16.20 1.00 16.65 1.00 17.94 1.00 19.95	B B B B B	C O N C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10263 10264 10265 10266 10267 10268 10269 10270		ASP ASP ASP THR THR THR THR	556 556 556 557 557 557	105. 355 105. 327 106. 862 106. 046 108. 039 108. 443 109. 923 110. 687	45. 429 45. 751 43. 202 42. 962 43. 762 44. 132 43. 826 44. 589	51. 583 49. 415 52. 597 53. 480 52. 847 54. 200 54. 396 53. 454	1.00 22.16 1.00 21.01 1.00 16.87 1.00 15.15 1.00 17.93 1.00 18.07 1.00 18.59	B B B B	0 0 0 0 N C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10271 10272 10273 10274 10275 10276 10277		THR THR THR VAL VAL VAL VAL	557 557 557 558 558 558 558	110. 188 108. 203 108. 776 107. 348 107. 049 106. 483 106. 033	42. 358 45. 616 46. 151 46. 272 47. 682 48. 302 49. 733	53. 454 54. 157 54. 531 55. 479 53. 754 53. 964 52. 676 52. 940	1. 00 20. 98 1. 00 19. 55 1. 00 17. 89 1. 00 16. 94 1. 00 16. 56 1. 00 14. 93 1. 00 14. 99 1. 00 13. 18	B B B B B	0 C C O N C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10278 10279 10280 10281 10282 10283 10284	CG2 C O N CA CB CG	VAL VAL PHE PHE PHE PHE	558 558 558 559 559 559	107. 544 106. 058 105. 060 106. 348 105. 484 106. 303 105. 469	48. 247 47. 921 47. 211 48. 923 49. 269 49. 933 50. 504	51.568 55.109	1. 00 13. 02 1. 00 15. 99 1. 00 13. 36 1. 00 15. 43 1. 00 14. 56 1. 00 12. 72 1. 00 11. 04	B B B B B	C C O N C C C	
ATOM ATOM ATOM ATOM ATOM	10285 10286 10287 10288 10289	CD1 CD2 CE1 CE2 CZ	PHE PHE	559 559 559 559 559	105. 064 105. 056 104. 260 104. 251 103. 855	49.712 51.833 50.232 52.360 51.554	60. 347 59. 244 61. 356 60. 252 61. 307	1.00 10.65 1.00 12.10 1.00 8.83 1.00 10.43 1.00 8.93	B B B B	C C C C C	

			FIG. 4	- 211		(Continued)
ATOM 1029 ATOM 1030 ATOM 1031 ATOM 1032 ATOM 10333 ATOM 10333 ATOM 10334 ATOM 10333 ATOM 10335 ATOM 10336 ATOM 10337 ATOM 10337 ATOM 10337 ATOM 10337	O PHE O PHE O PHE O PHE O PHE O NARG O ARG O CB ARG O NE ARG O NH2 ARG O NH2 ARG O LEU O ASN O LEU O LEU O ASN O CB TRP CCB TRP CC	559 1 560 1 560 1 560 1 560 1 560 1 560 1 560 1 560 1 560 1 561 1 561 1 561 1 561 1 561 561 1 562 99 1 562 99 562 99 1 562 99 562 99 562 99 563 563 563 563 563 563 563 563 563 563	FIG. 4 04. 395 50. 23 04. 696 51. 25 03. 137 49. 90 02. 029 50. 74 01. 354 50. 11 02. 248 49. 98 01. 491 49. 42 02. 322 48. 48. 82 03. 126 48. 82 03. 203 50. 090 03. 887 47. 918 00. 962 50. 980 00. 661 50. 100 00. 403 52. 183 00. 962 53. 878 00. 694 53. 878 00. 901 55. 299 00. 275 52. 934 00. 901 55. 299 00. 275 52. 934 00. 901 55. 299 00. 275 52. 934 00. 901 55. 299 00. 275 52. 934 00. 901 51. 841 6. 462 51. 267 6. 924 49. 823 7. 566 49. 309 6. 582 49. 157 4. 818 51. 139 4. 712 50. 793 3. 872 50. 936 2. 616 50. 281 1. 770 50. 132 0. 365 49. 719 9. 623 48. 721 4. 818 51. 139 4. 712 50. 793 3. 872 50. 936 2. 616 50. 281 1. 770 50. 132 0. 365 49. 719 9. 623 48. 721 3. 872 50. 936 2. 616 50. 281 1. 770 50. 132 0. 365 49. 719 9. 623 48. 721 3. 872 50. 936 2. 616 50. 281 1. 770 50. 132 0. 365 49. 719 9. 623 48. 721 3. 872 50. 936 2. 616 50. 281 1. 770 50. 132 48. 684 49. 959 48. 684 49. 959 48. 684 49. 959 48. 684 49. 959 48. 684 49. 973 48. 684 49. 980 48. 684 49. 973 48. 684 49. 980 48. 161 46. 186 585 46. 973 48. 037 48. 037	0 56. 592 1. 5 56. 000 1. 7 56. 865 1. 4 56. 421 1. 7 55. 185 1. 8 53. 954 1. 1 52. 755 1. 6 51. 999 1. 8 51. 002 1. 9 50. 614 1. 9 57. 486 1. 9 58. 291 1. 5 57. 483 1. 5 58. 392 1. 6 60. 189 1. 6 60. 189 1. 6 60. 698 1. 6 61. 310 1. 6 60. 698 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61. 310 1. 6 61.	00 10. 22 00 12. 59 00 12. 69 00 12. 69 00 15. 06 00 14. 07 00 14. 26 00 15. 38 00 15. 38 00 15. 38 00 15. 26 00 15. 26 00 15. 35 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 58 00 15. 64 00 15. 65 00 16. 15 00 16. 15 00 16. 15 00 16. 15 00 17. 65 00 16. 15 00 18. 07 00 18. 92	Continued) B C B O N C C C C C C C C C C C C C C C C C C
		SUBST	TITUTE SHEET	(RULE 26)		

										(((((((((((((((((((((((((((((((((((((((
					FIC	G. 4-	2 1 2			(Continued)
ATOM	10339	CA	THR	565	95.817	48. 259	60. 159	1.00 17.29	В	С
ATOM	10340	CB	THR	565	96.626	49. 551	60. 294	1.00 17.13	В	č
ATOM	10341		THR	565	97.677	49. 570	59. 330	1.00 20.36	В	ő
ATOM	10342		THR	565	97. 238	49.636	61.676	1.00 18.23	В	č
ATOM	10343	C	THR	565	94. 665	48. 355	61.157	1.00 15.84	B	č
ATOM	10344	Õ	THR	565	94. 738	47. 804	62. 249	1.00 14.07	В	ŏ
ATOM	10344	N	TYR	566	93. 605	49.061	60. 781	1.00 15.76	В	N N
ATOM	10346	CA	TYR	566	92. 455	49. 204	61.664	1.00 13.70	В	
ATOM	10347	CB	TYR	566	91. 543	50. 335	61. 177	1.00 17.74	В	C C C C C C
ATOM	10341	CG	TYR	566	90.067	50. 039	61.311	1.00 13.01	В	C
ATOM	10346		TYR	566	89. 303	49. 688	60. 195	1.00 17.40	В	C
ATOM	10345		TYR	566	87. 947	49. 390	60.310	1.00 17.77	B	C
ATOM	10350		TYR	566	89. 432	50.086	62.556	1.00 13.12	В	C
ATOM	10351		TYR	566		49. 789				C
ATOM	10352	CZ	TYR		88.073		62. 682	1.00 17.35	В	C
ATOM	10353	OH	TYR	566	87.340	49.441	61.550	1.00 17.10	В	0
ATOM		С		566 566	86.005	49.137	61.662	1.00 17.63	В	0
ATOM	10355 10356		TYR	566 566	91.667	47. 899	61.777	1.00 19.12	В	C
		0 N	TYR	566 567	91. 249	47.517	62.871	1.00 20.12	В	0 N
ATOM	10357	N	LEU	567 ·	91.481	47. 211	60.654	1.00 19.08	В	N
ATOM ATOM	10358	CA	LEU	567 567	90. 735	45. 959	60.648	1.00 19.66	В	C
	10359	CB	LEU	567	90.606	45. 419	59. 223	1.00 18.00	В	L O
ATOM	10360		LEU	567 .	89. 728	46. 252	58. 284	1.00 18.48	В	C C C C C
ATOM	10361		LEU	567 567	89. 735	45. 628	56. 889	1.00 19.22	В	Ü
ATOM	10362		LEU	567	88. 310	46. 325	58. 835	1.00 15.78	В	C
ATOM	10363	C	LEU	567 567	91: 355	44. 898	61.544	1.00 20.80	В	
ATOM	10364	0	LEU	567 568	90.645	44. 102	62. 157	1.00 23.88	В	0
ATOM	10365	N	ALA	568	92.677	44. 883	61.628	1.00 19.62	В	N
ATOM	10366	CA	ALA	568	93. 347	43.898	62.466	1.00 20.08	В	C
ATOM	10367	CB	ALA	568	94. 746	43.601	61.907	1.00 18.06	В	C
ATOM	10368	C	ALA	568	93. 451	44.362	63. 924	1.00 20.52	В	C
ATOM	10369	0	ALA	568	93. 319	43.569	64. 849	1.00 20.37	В	0
ATOM	10370	N	SER	569	93. 674	45.653	64. 128	1.00 20.79	В	N
ATOM	10371	CA	SER	569	93. 827	46. 182	65. 474	1.00 21.75	В	C
ATOM	10372	CB	SER	569	94. 520	47. 545	65. 401	1.00 21.85	В	C
ATOM	10373	0G	SER	569	94. 546	48. 188	66.657	1.00 22.64	В	0
ATOM	10374	C	SER	569	92. 525	46. 297	66. 267	1.00 22.83	В	Ç
ATOM	10375	0	SER	569	92. 505	46.029	67.470	1.00 22.38	В	0
ATOM	10376	N	THR	570	91.444	46.679	65. 589	1.00 22.26	В	N
ATOM	10377	CA	THR	570	90.153	46.862	66. 232	1.00 21.45	В	C
ATOM	10378	CB	THR	570	89. 512	48. 191	65.797	1.00 19.91	В	C
ATOM	10379		THR	570	90.349	49. 285	66.188	1.00 21.12	В	0
ATOM	10380		THR	570	88. 143	48.351	66.430	1.00 17.96	В	С
ATOM	10381	C	THR	570	89. 132	45. 751	65.974	1.00 24.43	В	C
ATOM	10382	0	THR	570	88. 453	45.301	66.894	1.00 27.79	В	0
ATOM	10383	N	GLU	571	89.001	45.317	64.727	1.00 23.34	В	N
ATOM	10384	CA	GLU	571	88. 030	44. 280	64.415	1.00 21.95	В	C
ATOM	10385	CB	GLU	571	87. 499	44. 481	62.998	1.00 22.83	В	C
ATOM	10386	CG	GLU	571	87.004	45.888	62.709	1.00 24.63	В	C
ATOM	10387	CD	GLU	571	85.957	46. 357	63.696	1.00 25.17	В	С

ATOM 10388 OE1 GLU 571 85.236 45.509 64.258 1.00 28.12 B O ATOM 10389 OE2 GLU 571 85.834 47.580 63.897 1.00 26.28 B O ATOM 10390 C GLU 571 88.606 42.874 64.554 1.00 21.35 B C ATOM 10391 O GLU 571 87.903 41.887 64.362 1.00 19.91 B O ATOM 10392 N ASN 572 89.887 42.784 64.894 1.00 22.55 B N ATOM 10393 CA ASN 572 90.539 41.491 65.043 1.00 21.58 B C ATOM 10394 CB ASN 572 89.998 40.744 66.255 1.00 23.76 B C ATOM 10395 CG ASN 572 90.523 41.303 67.552 1.00 27.80 B C	
ATOM 10391 O GLU 571 87.903 41.887 64.362 1.00 19.91 B O ATOM 10392 N ASN 572 89.887 42.784 64.894 1.00 22.55 B N ATOM 10393 CA ASN 572 90.539 41.491 65.043 1.00 21.58 B C ATOM 10394 CB ASN 572 89.998 40.744 66.255 1.00 23.76 B C	
ATOM 10393 CA ASN 572 90.539 41.491 65.043 1.00 21.58 B C ATOM 10394 CB ASN 572 89.998 40.744 66.255 1.00 23.76 B C	·
ATUM 10395 CG ASN 572 90.523 41.303 67.552 1.00 27.80 R C	
ATOM 10396 OD1 ASN 572 90.053 42.335 68.035 1.00 30.34 B 0	
ATOM 10397 ND2 ASN 572 91.522 40.634 68.121 1.00 30.31 B N ATOM 10398 C ASN 572 90.347 40.639 63.806 1.00 21.12 B C ATOM 10399 O ASN 572 90.112 39.436 63.903 1.00 20.16 B O	
ATOM 10400 N ILE 573 90.445 41.280 62.645 1.00 19.59 B N ATOM 10401 CA ILE 573 90.311 40.604 61.365 1.00 18.06 B C	
ATOM 10402 CB ILE 573 89.509 41.456 60.382 1.00 18.14 B C ATOM 10403 CG2 ILE 573 89.371 40.735 59.057 1.00 18.53 B C ATOM 10404 CG1 ILE 573 88.143 41.778 60.970 1.00 19.49 B C	
ATOM 10404 CG1 ILE 573 88.143 41.778 60.970 1.00 19.49 B C ATOM 10405 CD1 ILE 573 87.336 42.735 60.131 1.00 20.04 B C ATOM 10406 C ILE 573 91.706 40.425 60.777 1.00 18.47 B C	
ATOM 10407 0 ILE 573 92.480 41.376 60.739 1.00 19.08 B 0 ATOM 10408 N ILE 574 92.038 39.216 60.337 1.00 17.57 B N	
ATOM 10409 CA ILE 574 93.340 38.978 59.724 1.00 18.02 B C ATOM 10410 CB ILE 574 93.724 37.494 59.740 1.00 19.09 B C ATOM 10411 CG2 ILE 574 94.950 37.280 58.870 1.00 20.13 B C	
ATOM 10412 CG1 ILE 574 94.004 37.031 61.172 1.00 21.02 B C ATOM 10413 CD1 ILE 574 94.330 35.553 61.282 1.00 20.47 B C	
ATOM 10414 C ILE 574 93. 298 39. 423 58. 265 1. 00 17. 84 B C ATOM 10415 O ILE 574 92. 444 38. 981 57. 500 1. 00 19. 48 B O ATOM 10416 N VAL 575 94. 217 40. 296 57. 876 1. 00 17. 13 B N	
ATOM 10417 CA VAL 575 94.254 40.777 56.498 1.00 16.42 B C ATOM 10418 CB VAL 575 94.354 42.308 56.430 1.00 16.55 B C	
ATOM 10419 CG1 VAL 575 94. 271 42. 753 54. 985 1. 00 16. 06 B C ATOM 10420 CG2 VAL 575 93. 242 42. 948 57. 261 1. 00 15. 54 B C	
ATOM 10421 C VAL 575 95.452 40.187 55.786 1.00 16.02 B C ATOM 10422 O VAL 575 96.592 40.488 56.124 1.00 16.68 B O ATOM 10423 N ALA 576 95.186 39.344 54.797 1.00 16.21 B N	
ATOM 10424 CA ALA 576 96. 246 38. 683 54. 056 1. 00 15. 22 B C ATOM 10425 CB ALA 576 96. 062 37. 176 54. 127 1. 00 12. 38 B C	
ATOM 10426 C ALA 576 96.330 39.117 52.601 1.00 15.92 B C ATOM 10427 O ALA 576 95.397 39.710 52.046 1.00 16.20 B O ATOM 10428 N SER 577 97.470 38.811 51.996 1.00 14.35 B N	
ATOM 10429 CA SER 577 97.722 39.123 50.606 1.00 13.57 B C ATOM 10430 CB SER 577 98.368 40.495 50.474 1.00 13.58 B C	
ATOM 10431 OG SER 577 97. 456 41. 504 50. 866 1. 00 16. 22 B O ATOM 10432 C SER 577 98. 642 38. 045 50. 069 1. 00 13. 24 B C ATOM 10433 O SER 577 99. 497 37. 522 50. 788 1. 00 13. 05 B O	
ATOM 10434 N PHE 578 98.462 37.712 48.800 1.00 11.98 B N ATOM 10435 CA PHE 578 99.262 36.676 48.183 1.00 11.24 B C ATOM 10436 CB PHE 578 98.418 35.407 48.079 1.00 11.42 B C	

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		٠.		-	FI	G. 4-	214		•	(Continued)
ATOM ATOM	10437 10438	CG CD1	PHE ·	578 578	99. 136 100. 196	34. 232	47. 481 48. 152	1.00 10.60 1.00 10.29	B B	C C C C C
ATOM	10439	CD2	PHE	578	98. 697		46. 280	1.00 10.25	В	č
ATOM	10440		PHE	578	100.805		47.640	1.00 11.15	В	Ç
ATOM	10441		PHE	578 578	99. 297		45.762	1.00 11.72	В	C
ATOM ATOM	10442 10443	CZ C	PHE PHE	578 578	100. 354 99. 746		46. 446 46. 805	1.00 10.87 1.00 10.56	B B	C
ATOM	10444	Ö	PHE	578	99.002		46.039	1.00 10.36	В	Ö
ATOM	10445	Ň	ASP	579	101.005		46.516	1.00 11.14	В	Ň
ATOM	10446	CA	ASP	579	101.617		45. 227	1.00 9.94	·B	Ċ
ATOM	10447	CB	ASP	579	103.008	37.682	45.401	1.00 9.15	В	C C C
ATOM	10448	CG	ASP	579	102.957	39.090	45.954	1.00 13.00	В	
ATOM	10449		ASP	579	102.053		45.532	1.00 14.87	В	0
ATOM ATOM	10450 10451	C	ASP ASP	579 579	103. 816 101. 734		46.796	1.00 11.19 1.00 11.60	В	0
ATOM	10451	0	ASP	579	101. 734		44. 488 44. 753	1.00 11.00	B B	C 0
ATOM	10453	Ň	GLY	580	100. 809	35. 510	43. 570	1.00 10.77	В	N
ATOM	10454	CA	GLY	580	100. 838		42. 815	1.00 11.96	B	Ċ
ATOM	10455	C	GLY	580	101.458	34.470	41.450	1.00 13.34	В	C
ATOM	10456	0	GLY	580	102. 269	35.376	41.227	1.00 12.96	В	0
ATOM	10457	N	ARG .	581	101.080	33.611	40. 521	1.00 14.18	В	N
ATOM	10458	CA	ARG	581	101.615	33.714	39. 187	1.00 15.34	В	C
ATOM ATOM	10459 10460	CB CG	ARG ARG	581	101.085	32. 570	38. 338	1.00 13.67	В	C
ATOM	10460	CD	ARG	581 581	101.809 101.172	31.283 30.076	38.666 38.023	1.00 15.30 1.00 14.62	B B	C C
ATOM	10462	NE	ARG	581	99. 980	29.652	38. 740	1.00 14.02	В	N N
ATOM	10463	CZ	ARG	581	99. 186	28. 672	38. 330	1.00 13.69	В	Č
ATOM	10464	NH1	ARG	581	99.467	28.024	37. 207	1.00 13.99	B	Ň
ATOM	10465		ARG	581	98.112	28.348	39.036	1.00 12.41	В	N
ATOM	10466	C	ARG	581	101. 237	35.069	38.624	1.00 17.21	В	C
ATOM	10467	0	ARG	581	100.175	35. 615	38. 934	1.00 17.96	В	0
ATOM	10468	N	GLY	582	102. 128	35.628		1.00 18.14	B	N
ATOM ATOM	10469 10470	CA C	GLY GLY	582 582	101.868 102.454	36. 933 37. 998	37. 258 38. 159	1.00 17.73 1.00 16.81	B B	C
ATOM	10470	Ö	GLY	582	102. 454	39. 151	37. 754	1.00 10.01	В	0
ATOM	10472	Ň	SER	583	102. 835	37. 625	39. 378	1.00 15.90	В	N
ATOM	10473	CA	SER	583	103. 423	38. 588	40. 309	1.00 16.60	B	Č
ATOM	10474	CB	SER	583	103. 437	38.024	41.730	1.00 17.47	B	Č
ATOM	10475	0G	SER	583	104. 229	36.856	41.811	1.00 21.54	В	0
ATOM	10476	C	SER	583	104.841	38. 901	39. 841	1.00 15.56	В	С
ATOM	10477	0	SER	583	105. 389	38.176		1.00 17.79	В	0
ATOM ATOM	10478	N CA	GLY	584	105.441		. 40. 359	1.00 14.64	. B	N
ATOM	10479 10480	CA	GLY GLY	584 584	106. 776 107. 969	40. 334 40. 158	39. 908 40. 831	1.00 13.05 1.00 12.28	B B	C
ATOM	10481	0	GLY	, 584	107. 851	39.648	40. 631	1.00 12.28	В	C 0
ATOM	10482	N	TYR	585	109. 129	40. 583	40. 325	1.00 11.78	• В	N
ATOM	10483	CA	TYR	585	110.412	40.536	41.034	1.00 12.19	В	Ċ
ATOM	10484	CB	TYR	585	110.335	41.383	42. 304	1.00 11.93	B	C
ATOM	10485	CG	TYR	585	109.704	42.719	42.047	1.00 12.41	В	C

					FIC	G. 4-	215	·	-	(Continued)
ATOM ATOM ATOM	10486 10487 10488 10489	CE1 CD2 CE2	TYR TYR TYR TYR	585 585 585 585	110. 370 109. 756 108. 408 107. 783	43. 694 44. 891 42. 983 44. 179	41. 297 40. 979 42. 478 42. 167	1.00 12.30 1.00 12.43 1.00 10.95 1.00 12.28	B B B	C C C
ATOM ATOM ATOM ATOM ATOM	10490 10491 10492 10493 10494	CZ OH C O N	TYR TYR TYR TYR GLN	585 585 585 585 586	108. 459 107. 831 110. 883 111. 673 110. 413	45. 126 46. 306 39. 141 38. 979 38. 144	41. 418 41. 109 41. 394 42. 319 40. 655	1.00 13.31 1.00 14.33 1.00 12.01 1.00 13.01 1.00 11.45	B B B B	C O C O N
ATOM ATOM ATOM ATOM ATOM	10495 10496 10497 10498	CA CB CG CD	GLN GLN GLN GLN	586 586 586 586	110. 787 109. 639 109. 178 107. 749	36. 763 36. 071 36. 854 36. 533	40. 906 41. 641 42. 867 43. 295	1.00 11.62 1.00 10.30 1.00 14.38 1.00 15.38	B B B	C C C
ATOM ATOM ATOM ATOM	10499 10500 10501 10502 10503	NE2 C O N	GLN GLN GLN GLN GLY	586 586 586 586 587	107. 468 106. 835 111. 118 111. 173 111. 336	35. 452 37. 478 36. 023 34. 786 36. 778	43. 816 43. 060 39. 602 39. 574 38. 525	1.00 12.14 1.00 15.36 1.00 12.85 1.00 13.97 1.00 11.70	B B B B	0 N C O N
ATOM ATOM ATOM ATOM ATOM	10504 10505 10506 10507 10508	CA C O N CA	GLY GLY GLY ASP ASP	587 587 587 588 588	111. 641 110. 405 109. 302 110. 595 109. 500	36. 168 35. 960 35. 786 35. 949 35. 776	37. 242 36. 373 36. 884 35. 054 34. 105	1.00 11.61 1.00 14.10 1.00 13.91 1.00 16.19 1.00 17.70	B B B B	C C O N C
ATOM ATOM ATOM ATOM ATOM	10509 10510 10511 10512 10513	CB CG OD1	ASP ASP ASP ASP	588 588 588 588	110.002 110.708 110.236 111.738	35. 993 37. 312 38. 335 37. 327	32. 680 32. 505 33. 040 31. 809	1.00 18.98 1.00 20.57 1.00 23.28 1.00 23.25	В В В	C C O O
ATOM ATOM ATOM ATOM	10514 10515 10516 10517	O N CA CB	ASP LYS LYS LYS	588 588 589 589 589	108. 723 107. 608 109. 294 108. 559 109. 383	34. 454 34. 389 33. 397 32. 143 31. 030	34. 139 33. 635 34. 697 34. 734 35. 372	1.00 17.46 1.00 16.74 1.00 18.02 1.00 20.00 1.00 22.21	B B B B	C O N C
ATOM ATOM ATOM ATOM ATOM	10518 10519 10520 10521 10522	CG CD CE NZ C	LYS LYS LYS LYS LYS	589 589 589 589 589	108. 633 109. 526 108. 753 109. 605	29. 710 28. 579 27. 273 26. 232	35. 443 35. 940 36. 111 36. 771	1.00 27.16 1.00 32.47 1.00 33.79 1.00 35.98	В В В В	C C C N
ATOM ATOM ATOM ATOM	10523 10524 10525 10526	O N CA CB	LYS ILE ILE ILE	589 590 590 590	107. 290 106. 244 107. 384 106. 237 106. 681	32. 362 31. 781 33. 212 33. 523 33. 901	35. 536 35. 242 36. 552 37. 379 38. 814	1.00 20.94 1.00 23.79 1.00 18.06 1.00 14.07 1.00 11.33	B B B B	C O N C C
ATOM ATOM ATOM ATOM	10527 10528 10529 10530	CG1 CD1 C	ILE	590 590 590 590	105. 585 107. 057 107. 750 105. 461	34. 654 32. 635 32. 888 34. 682	39. 538 39. 585 40. 897 36. 753	1. 00 9. 61 1. 00 10. 89 1. 00 7. 05 1. 00 15. 70	В В В В	C C C
ATOM ATOM ATOM ATOM	10531 10532 10533 10534	O N CA CB	ILE MET MET MET	590 591 591 591	104. 254 106. 159 105. 506 106. 512	34. 583 35. 774 36. 948 38. 088	36. 511 36. 465 35. 907 35. 759	1. 00 16. 31 1. 00 15. 00 1. 00 14. 79 1. 00 14. 22	B B B	O N C C

					FIG. 4-216	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10535 10536 10537 10538 10539 10540 10541 10542 10543 10544 10545 10550 10551 10552 10553 10555 10556 10557 10558 10559 10560	ND1 CE1 NE2 C O N CA CB C CO N CA CB CG2 CG1 CD1	ILE ILE	591 591 591 591 592 592 592 592 592 592 593 593 593 594 594 594 594 594	105. 854 39. 452 35. 581 1.00 18. 55 B 107. 027 40. 830 35. 526 1.00 17. 84 B 107. 813 40. 502 33. 933 1.00 16. 39 B 104. 788 36. 699 34. 582 1.00 14. 86 B 103. 643 37. 113 34. 418 1.00 14. 45 B 105. 451 36. 022 33. 647 1.00 14. 66 B 104. 863 35. 725 32. 343 1.00 14. 33 B 105. 962 35. 424 31. 332 1.00 15. 14 B 106. 753 36. 626 30. 922 1.00 17. 56 B 106. 626 37. 933 31. 252 1.00 17. 20 B 107. 810 36. 555 30. 041 1.00 17. 84 B 108. 300 37. 765 29. 845 1.00 16. 59 B 107. 598 38. 620 30. 567 1.00 16. 88 B 103. 224 34. 274 31. 344 1.00 15. 86 B 102. 775 32. 810 33. 615 1.00 15. 66 B 100. 647	S C C O N C C C C N C N C O N C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10559 10560 10561 10562 10563 10564 10565 10566 10567 10570 10571 10572 10573 10574 10575 10576	CG1 CD1 C O N CA CB CG OD1 ND2 C O N CA CB CG CB CC N N CA CB	ILE ILE ILE ASN ASN ASN ASN ASN ASN ASN ARG ARG ARG ARG ARG	594 594 594 595 595 595 595 596 596 596 596 596	100. 006 36. 915 34. 321 1. 00 16. 86 B 100. 533 37. 882 33. 274 1. 00 16. 67 B 99. 748 35. 689 31. 413 1. 00 17. 96 B 98. 884 36. 525 31. 160 1. 00 19. 03 B 100. 718 35. 385 30. 558 1. 00 17. 93 B 100. 802 36. 050 29. 263 1. 00 19. 09 B 102. 140 35. 737 28. 592 1. 00 19. 22 B 102. 291 36. 441 27. 260 1. 00 19. 91 B 102. 320 37. 668 27. 198 1. 00 19. 01 B 102. 377 35. 667 26. 184 1. 00 19. 95 B 99. 659 35. 641 28. 330 1. 00 19. 95 B 99. 456 34. 460 28. 076 1. 00 19. 31 B 98. 933 36. 630 27. 814 1. 00 19. 66 B 97. 799 36. 406 26. 911 1. 00 20. 07 B 98. 212 35. 588 25. 677 1. 00 17. 78 B 99. 655	C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10577 10578 10579 10580 10581 10582 10583	CZ NH1 NH2 C O N CA	ARG ARG ARG ARG ARG ARG ARG	596 596 596 596 596 597	98. 102 35. 816 21. 795 1. 00 19. 85 B 98. 671 37. 005 21. 640 1. 00 21. 47 B 97. 060 35. 486 21. 045 1. 00 18. 12 B 96. 692 35. 655 27. 632 1. 00 21. 03 B 95. 731 35. 213 27. 005 1. 00 22. 67 B 96. 811 35. 529 28. 948 1. 00 20. 90 B 95. 831 34. 770 29. 714 1. 00 20. 85 B	C N N C O N C

		٠			FI	G. 4	- 217			(Cont	tinued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10584 10585 10586 10587 10588 10589 10590 10591 10592	NH2 C	ARG ARG ARG ARG ARG ARG ARG	597 597 597 597 597 597 597	96. 437 95. 850 95. 913 95. 006 94. 776 95. 386 93. 933 95. 292	32. 257 32. 520 31. 660 31. 792 32. 748 30. 974 35. 429	29. 300 27. 810 27. 059 25. 759 25. 075 25. 145 30. 976	1. 00 23. 88 1. 00 31. 40 1. 00 34. 67 1. 00 35. 49 1. 00 35. 98 1. 00 35. 20 1. 00 39. 12 1. 00 18. 83	B B B B B	C C C N C N C	
ATOM ATOM ATOM ATOM ATOM ATOM	10593 10594 10595 10596 10597 10598 10599	CD2 C	ARG LEU LEU LEU LEU LEU LEU	597 598 598 598 598 598 598	94. 981 95. 175 94. 678 94. 482 95. 523 96. 939 95. 361 93. 369	36. 751 37. 477 38. 959 39. 990 39. 473 41. 267 36. 870	30. 964 32. 125 31. 769 32. 248 32. 106 31. 466 32. 642	1.00 17.54 1.00 16.66 1.00 15.71 1.00 12.95 1.00 12.69 1.00 11.40 1.00 9.68 1.00 17.19	B B B B B	O N C C C C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10600 10601 10602 10603 10604 10605 10606 10607	O N CA C O N CA CB	LEU GLY GLY GLY THR THR THR	598 599 599 599 599 600 600	92. 533 93. 207 91. 997 91. 987 90. 921 93. 164 93. 247 93. 823	36. 398 36. 864 36. 328 34. 824 34. 221 34. 213 32. 775 32. 091	31. 863 33. 961 34. 547 34. 735 34. 843 34. 786 34. 972 33. 722	1.00 17.25 1.00 16.06 1.00 16.44 1.00 17.37 1.00 17.28 1.00 17.73 1.00 17.95 1.00 18.93	B B B B B B B	0 N C C O N C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10608 10609 10610 10611 10612 10613 10614 10615		THR THR THR PHE PHE PHE PHE	600 600 600 600 601 601 601	95. 185 93. 000 94. 087 93. 574 95. 382 96. 279 97. 686 97. 757	32. 495 32. 463 32. 384 32. 285 32. 177 31. 768 31. 542 30. 452	33. 530 32. 491 36. 183 37. 295 35. 971 37. 048 36. 494 35. 475	1.00 17.74 1.00 17.48 1.00 19.93 1.00 21.69 1.00 21.11 1.00 21.56 1.00 20.77 1.00 21.75	B B B B B	0 C C O N C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10616 10617 10618 10619 10620 10621 10622 10623	CD1 CD2 CE1	PHE PHE PHE PHE PHE PHE PHE	601 601 601 601 601 601 601 602	98. 676 96. 896 98. 731 96. 949 97. 868 96. 346 96. 437 96. 312	30. 513 29. 366 29. 502 28. 356 28. 427 32. 710 32. 247 34. 018	34. 439 35. 539 33. 474 34. 581 33. 547 38. 244 39. 386	1.00 23.50 1.00 21.83 1.00 24.75 1.00 22.61 1.00 20.03 1.00 21.61 1.00 23.03	B B B B B	C C C C C C C C C C C C C C C C C C C	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10624 10625 10626 10627 10628 10629 10630 10631 10632	CA CB CG CD OE1 OE2 C	GLU GLU GLU GLU GLU	602	96. 374 96. 505 95. 193 94. 857 94. 930 94. 505 95. 111 95. 170 93. 979	34. 976 36. 422 37. 072 36. 847 35. 696 37. 830 34. 838 34. 953 34. 584	37. 997 39. 097 38. 581 38. 135 36. 661 36. 184 35. 981 39. 952 41. 179 39. 296	1.00 20.14 1.00 19.30 1.00 16.90 1.00 17.16 1.00 17.31 1.00 18.92 1.00 16.38 1.00 18.97 1.00 18.54 1.00 19.02	B B B B B B	N C C C O O C O N	

					E 1.	7 4	0 1 0			(Continued)
					Г 1 (G. 4-	218			
ATOM	10633	CA	VAL	603	92.696	34.413	39. 984	1.00 21.62	В	C C
ATOM	10634	CB	VAL	603	91.513	34. 471	38. 999	1.00 21.51	В	C
ATOM	10635		VAL	603	90. 233	34.055	39. 701	1.00 19.24	В	C
ATOM	10636		VAL	603	91.380	35.876	38. 442	1.00 21.00	В	C
ATOM	10637	C	VAL	603	92. 643	33.073	40.716	1.00 22.35	В	C
MOTA	10638	0 M	VAL	603	92. 160	32. 989	41.848	1.00 21.06	В	0
MOTA	10639	N CA	GLU	604	93. 141	32. 031	40.059	1.00 22.98	В	N C
ATOM ATOM	10640 10641	CA CB	GLU GLU	604 604	93. 182 93. 721	30. 702 29. 681	40.656	1.00 26.04	В	C
ATOM	10642	CG	GLU	604	92. 956	29. 671	39. 645 38. 326	1.00 28.46 1.00 35.94	B B	C C
ATOM	10643	CD	GLU	604	93. 559	28. 742	37. 273	1.00 33.34	В	Č
ATOM	10644		GLU	604	93. 215	28. 911	36.076	1.00 40.17	В	Ö
ATOM	10645		GLU	604	94. 360	27.844	37. 637	1.00 41.61	В	Ŏ
ATOM	10646	C	GLU	604	94. 072	30. 705	41.905	1.00 24.63	B	č
ATOM	10647	Ō	GLU	604	93.657	30. 255	42.976	1.00 25.47	B	ŏ
ATOM	10648	N	ASP	605	95. 286	31.234	41.775	1.00 22.17	B	N
ATOM	10649	CA	ASP	605	96. 213	31.255	42.900	1.00 21.12	В	C
ATOM	10650	CB	ASP	605	97.568	31.827	42.463	1.00 23.09	В	C
ATOM	10651	CG	ASP	605	98. 263	30.958	41.414	1.00 24.43	В	C
ATOM	10652		ASP	605		29.774	41.266	1.00 26.59	В	0
ATOM	10653		ASP	605	99. 188	31.453	40.742	1.00 25.60	В	0
ATOM	10654	C	ASP	605	95. 712	31.967	44.159	1.00 19.42	В	C
ATOM	10655	0	ASP	605	96. 099	31.598	45. 260	1.00 19.67	В	0
ATOM	10656	N	GLN	606	94. 868	32. 983	44.014	1.00 17.23	В	N
ATOM	10657	CA	GLN	606	94. 337	33. 673	45.192	1.00 16.41	В	C
ATOM ATOM	10658 10659	CB CG	GLN	606	93. 576	34. 951	44. 795	1.00 17.09	В	C
ATOM	10660	CD	GLN GLN	606 606	94. 407	36.070	44.165	1.00 15.81	В	C
ATOM	10661		GLN	606	95. 332 94. 879	36. 748 37. 283	45. 162 46. 173	1.00 15.36	В	C
ATOM	10662		GLN	606	96. 637	36. 730	40.173	1.00 13.19 1.00 14.39	B B	0 · N
ATOM	10663	C	GLN	606	93. 360	32. 706	45. 878	1.00 14.33	В	C
ATOM	10664	ŏ	GLN	606	93. 337	32. 583	47. 102	1.00 14.30	В	0
ATOM	10665	Ň	ILE	607	92. 549	32.030	45.070	1.00 13.95	В	N
ATOM	10666	CA	ILE	607	91.584	31.076	45.583	1.00 13.95	B	Ç ·
ATOM	10667	CB	ILE	607	90.772	30.437	44.448	1.00 12.90	B	č
ATOM	10668		ILE	607	89. 925	29. 294	44.996	1.00 11.78	B	Č
ATOM	10669	CG1	ILE	607	89.909	31.504	43.773	1.00 12.90	В	Č
ATOM	10670		ILE	607	89. 162	31.016	42.560	1.00 11.00	В	C
ATOM	10671	C	ILE	607	92.330	29.985	46.318	1.00 15.04	В	C
ATOM	10672	0	ILE	607	92.008	29.670	47.462	1.00 15.40	В	0
ATOM	10673	N	GLU	608	93. 331	29.413	45.652	1.00 16.29	В	N
ATOM	10674	CA	GLU	608	94. 144	28. 359	46.246	1.00 18.48	В	C
ATOM	10675	CB	GLU	608	95. 180	27. 864	45. 235	1.00 18.74	В	C
ATOM	10676	CC	GLU	608	96.164	26.851	45. 792	1.00 22.43	В	C C C
ATOM	10677	CD	GLU	608	95.498	25. 557	46. 213	1.00 29.00	В	
ATOM	10678		GLU GLU	608	96.096	24. 817	47.032	1.00 32.52	В	0
ATOM ATOM	10679 10680	C	GLU	608 608	94. 382 94. 848	25. 274	45. 721	1.00 31.62	В	0
ATOM	10681	0	GLU	608	95.114	28. 889 28. 138	47. 501 48. 446	1.00 20.58 1.00 23.01	B B	C 0
111 Old	10001	U	OHO	000	JU. 114	40. 100	40.440	1.00 20.01	D	U

					FIG. 4-219	-	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10682 10683 10684 10685 10686 10687 10699 10691 10692 10693 10694 10695 10696 10697 10698 10699 10700 10701 10702 10703 10704 10705 10706 10707 10708 10709 10710 10711 10712 10713	CA CB C O N CA CB CC O N CA CC O N CA CC O N CA	ALA ALA ALA ALA ALA ARG ARG ARG GLN GLN GLN GLN PHE PHE	609 609 609 609 610 610 610 611 611 611 611 611 612 612 612 612 612	FIG. 4 - 219 95.150 30.183 47.506 1.00 19.99 95.811 30.789 48.646 1.00 21.28 96.269 32.196 48.310 1.00 19.81 94.826 30.819 49.797 1.00 21.63 95.152 30.426 50.915 1.00 21.88 93.618 31.286 49.516 1.00 23.07 92.580 31.358 50.535 1.00 25.56 91.317 31.963 49.957 1.00 25.38 92.300 29.952 51.024 1.00 26.13 92.256 29.694 52.223 1.00 25.97 92.119 29.044 50.073 1.00 28.12 91.838 27.647 50.374 1.00 28.88 91.886 26.826 49.087 1.00 27.27 91.518 25.372 49.260 1.00 28.40 91.547 24.668 47.925 1.00 30.54 90.501 25.152 47.028 1.00 33.73 90.628 25.223 45.706 1.00 36.39 91.764 24.848 45.129 1.00 36.39 91.764 24.848 45.129 1.00 38.00 89.615 25.645 44.956 1.00 37.15 92.826 27.082 51.391 1.00 29.24 92.446 26.330 52.287 1.00 30.51 94.092 27.452 51.260 1.00 30.51 94.092 27.452 51.260 1.00 30.75 96.491 27.029 51.532 1.00 29.62 96.738 25.866 50.581 1.00 31.27 98.183 25.741 50.150 1.00 32.19 99.097 25.778 50.979 1.00 32.20 98.400 25.578 48.848 1.00 31.86 95.109 27.691 53.524 1.00 31.36 95.441 27.095 54.545 1.00 32.39 94.740 28.969 53.533 1.00 31.39 94.705 29.717 54.784 1.00 30.50	B B B B B B B B B B B B B B B B B B B	(Continued) N C C C C O N C C C C C N C C C C N C C C C
ATOM ATOM ATOM	10714 10715 10716	CB CG CD1	PHE PHE PHE	613 613 613	94. 527 31. 217 54. 538 1. 00 30. 43 95. 651 31. 853 53. 775 1. 00 31. 06	B B	C
ATOM ATOM ATOM ATOM	10717 10718 10719 10720	CD2 CE1 CE2 CZ	PHE PHE PHE PHE	613 613 613 613	95. 385 32. 805 52. 796 1. 00 30. 25 98. 024 32. 156 53. 371 1. 00 32. 97 96. 419 33. 432 52. 109 1. 00 31. 17 97. 742 33. 109 52. 394 1. 00 32. 13	B B B B	C C C C
ATOM ATOM ATOM ATOM		C O N CA CB	PHE PHE SER SER SER	613 613 614 614	93. 531 29. 214 55. 607 1. 00 30. 36 93. 572 29. 216 56. 830 1. 00 28. 96 92. 478 28. 786 54. 923 1. 00 31. 88 91. 292 28. 286 55. 600 1. 00 34. 43 90. 141 28. 104 54. 607 1. 00 34. 30	B B B B	C O N C C
ATOM ATOM ATOM ATOM ATOM	10726 10727 10728 10729 10730	OG C O N CA	SER SER SER LYS LYS	614 614 614 615 615	90. 419 27. 055 53. 697 1. 00 34. 39 91. 609 26. 953 56. 264 1. 00 35. 74 90. 908 26. 519 57. 178 1. 00 37. 21 92. 670 26. 307 55. 797 1. 00 36. 52 93. 079 25. 030 56. 350 1. 00 37. 25	B B B B	0 C 0 N C

				FIG. 4-220	(Con	tinued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10731 10732 10733 10734 10735 10736 10737 10738 10739 10740 10741 10742 10743 10744 10745 10746 10747 10753 10753 10754 10755 10756 10757 10768 10761 10762 10763 10764 10765 10763 10764 10765 10766 10767 10768 10767 10778 10778 10778 10778 10779	CD2 CE1 CE2 CZ C O N CA CB CG1 CG2 C O N CA CB CG OD1 OD2 C O N CA	615 615 615 615 615 616 616 616 616 616	93. 781 24. 196 55. 283 1. 00 37. 94 B 92. 839 23. 516 54. 293 1. 00 40. 25 B 93. 595 23. 050 53. 053 1. 00 42. 18 B 94. 883 22. 317 53. 419 1. 00 42. 76 B 95. 776 22. 147 52. 237 1. 00 43. 07 B	CCCCNCONCCCSCCONCCONCCCCCCCCCONCCCCCONCCCCONCCCC	

					FIG. 4-221	(Continue	d)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	10780 10781 10782 10783 10784 10785 10786 10787 10788 10799 10791 10792 10793 10794 10795 10796 10797 10798 10799 10800 10801 10802 10803 10804 10805 10806 10807 10808 10809 10811 10812 10813 10814 10815 10818	NDC O N CA CB CC CD CO N CA CB CC CD CO N CA	ASN LYS LYS LYS LYS LYS ARG ARG ARG ARG ARG ARG ARG ARG ALE ILE ILE ALA ALA ALA ILE	621 621 622 622 622 622 622 622 623 623 623 623	84. 795 26. 940 57. 415 1. 00 38. 23 86. 630 25. 763 57. 972 1. 00 39. 37 85. 310 29. 639 56. 756 1. 00 29. 63 84. 887 29. 626 55. 604 1. 00 30. 93 84. 563 30. 007 57. 787 1. 00 28. 32 83. 195 30. 441 57. 573 1. 00 27. 00 82. 303 29. 986 58. 740 1. 00 29. 24 82. 062 28. 471 58. 738 1. 00 32. 47 81. 029 28. 002 59. 761 1. 00 33. 84 81. 527 28. 099 61. 197 1. 00 36. 73 83. 168 31. 957 57. 404 1. 00 25. 42 82. 145 32. 543 57. 047 1. 00 26. 19 84. 314 32. 583 57. 642 1. 00 21. 83 84. 436 34. 023 57. 515 1. 00 18. 89 84. 380 34. 664 58. 895 1. 00 16. 79 83. 122 34. 394 60. 991 1. 00 19. 29 83. 405 35. 632 61. 690 1. 00 19. 11 84. 207 35. 718 <	B O N C O C C C C C C C C C C C C C C C C	d)
ATOM ATOM	10819 10820	CB CG2	ILE ILE	626 626	88. 207 38. 397 48. 478 1. 00 19. 21 B 86. 883 38. 365 47. 742 1. 00 19. 01 B	C	
ATOM ATOM ATOM	10821 10822 10823	CD1 C	ILE ILE ILE	626 626 626	89. 348 38. 713 47. 511 1. 00 18. 94 B 89. 576 37. 642 46. 471 1. 00 20. 78 B 87. 850 40. 810 49. 003 1. 00 17. 46 B	C	
	10824 10825 10826	O N CA	ILE TRP TRP	626 627 627	86. 692 41. 116 48. 754 1. 00 18. 15 B 88. 878 41. 628 48. 781 1. 00 16. 65 B 88. 663 42. 938 48. 177 1. 00 15. 95 B	O N C	
ATOM ATOM	10827 10828	CB CG	TRP TRP	627 627	88. 215 43. 945 49. 231 1. 00 14. 07 89. 318 44. 713 49. 875 1. 00 12. 00 B	C C	

					-					
										(Continued)
					FIC	3. 4 -	222			
ATOM	10829	CD2	TRP	627	89.641	46.084	49.646	1.00 11.41	В	C
ATOM	10830	CE2	TRP	627	90.725	46.410	50.500	1.00 10.99	В	C
ATOM	10831	CE3	TRP	627	89.121	47.074	48.806	1.00 9.75	В	C
ATOM	10832		TRP	627	90.198	44. 267	50.826	1.00 14.55	В	С
ATOM	10833		TRP	627	91.046	45. 283	51.208	1.00 10.25	В	N
ATOM	10834		TRP	627	91.289	47. 681	50. 536	1.00 9.06	B	C
ATOM	10835		TRP	627	89.685	48. 340	48.844	1.00 9.47	В	Č
ATOM	10836		TRP	627	90. 755	48. 632	49.702	1.00 8.43	В	C
ATOM	10837	C	TRP	627	89. 881	43. 489	47. 433	1.00 17.27	B	Č
ATOM	10838	ŏ	TRP	627	91.027	43. 146	47. 732	1.00 16.96	B	0
ATOM	10839	Ň	GLY	628	89.613	44. 351	46. 459	1.00 16.52	B	N
ATOM	10840	CA	GLY	628	90.672	44. 947	45.675	1.00 16.52	B	Ĉ
ATOM	10841	C	GLY	628	90.186	46. 198	44. 975	1.00 17.44	B	č
ATOM	10842	ŏ	GLY	628	88.977	46. 441	44. 887	1.00 17.88	B	Ŏ -
ATOM	10843	Ň	TRP	629	91.132	46. 989	44. 479	1.00 15.93	B	Ň
ATOM	10844	CA	TRP	629	90.841	48. 235	43. 781	1.00 15.93	B	Ċ
ATOM	10845	CB	TRP	629	91.480	49. 395	44. 552	1.00 13.57	В	č
ATOM	10846	CG	TRP	629	90.867	50. 763	44. 341	1.00 14.96	B	č
ATOM		· CD2		629	90.389	51.656	45. 360	1.00 13.15	B	č
ATOM	10848		TRP	629	89.944	52. 830	44.712	1.00 13.17	B	Č
ATOM	10849		TRP	629	90. 296	51.577	46. 758	1.00 14.07	B	Č
ATOM	10850		TRP	629	90.694	51.419	43. 149	1.00 14.45	B	Č
ATOM	10851		TRP	629	90. 141	52.657	43.366	1.00 12.77	B	N
ATOM	10852		TRP	629	89.411	53. 921	45.414	1.00 13.59	B	Ċ
ATOM	10853		TRP	629	89.767	52.660	47.461	1.00 14.81	B	Č
ATOM	10854		TRP	629	89.330	53.820	46.782	1.00 15.16	B	Č
ATOM	10855	C	TRP	629	91.481	48.074	42.399	1.00 17.34	B	Č
ATOM	10856		TRP	629	92.571	47.517	42.285	1.00 18.55	B	0
ATOM	10857	N	SER		90.802	48.538	41.354	1.00 17.70	В	N
ATOM	10858	CA	SER	630	91.309	48.430	39. 982	1.00 17.70	В	C
ATOM	10859	CB	SER	630	92.649	49. 144	39.846	1.00 18.19	В	C
ATOM	10860	0G	SER	630	92.574	50.437	40.404	1.00 24.67	В	0
ATOM	10861	C	SER	630	91.477	46.977	39.563	1.00 17.40	В	C
ATOM	10862	0	SER	630	90.501	46. 235	39.469	1.00 18.69	В	0
ATOM	10863	N	TYR	631	92.712	46.565	39.304	1.00 16.34	В	N
ATOM	10864	CA	TYR	631	92.951	45.192	38.904	1.00 15.96	В	С
ATOM	10865	CB	TYR	631	94.430	44.973	38.579	1.00 15.36	В	C
ATOM	10866	CG	TYR	631	94.689	43.709	37.779	1.00 15.93	В	
ATOM	10867	CD1	TYR	631	94.626	42.450	38.380	1.00 15.38	В	C C C
ATOM	10868	CE1	TYR	631	94.830	41.287	37.634	1.00 16.25	В	C
ATOM	10869		TYR	631	94.961	43.773	36.409	1.00 15.67	В	C
ATOM	10870	CE2	TYR	631	95.160	42.620	35.655	1.00 13.59	В	C
ATOM	10871	CZ	TYR	631	95.092	41.384	36.270	1.00 15.96	В	C
ATOM	10872	0H	TYR	631	95.264	40.243	35. 525	1.00 14.59	В	0
ATOM	10873	C	TYR	631	92.499	44. 286	40.049	1.00 15.68	В	C
ATOM	10874	0	TYR	631	91.949	43.213	39.824	1.00 16.42	В	0
ATOM	10875	N	GLY	632	92.723	44.729	41.281	1.00 15.56	В	N
ATOM	10876	CA	GLY	632	92.292	43.950	42.429	1.00 14.43	В	С
ATOM	10877	C	GLY	632	90.777	43.807	42.398	1.00 13.07	В	С

					FΙ	G. 4	- 223			(Co	ontinued)
ATOM ATOM	10878 10879	0 N	GLY GLY	632	90. 23	9 42.77	1 42.777	1.00 12.0			
ATOM	10880	CA	GLY	633 633	90. 08 88. 63						
ATOM	10881	C	GLY	633	88. 27						
ATOM	10882	0	GLY	633	87. 33				26 B	0	
ATOM	10883	N	TYR	634	89.03						
ATOM ATOM	10884 10885	CA CB	TYR TYR	634	88. 82)9 B		
ATOM	10886	CG	TYR	634 634	89. 86 89. 81			1.00 7.3 1.00 8.0			
ATOM	10887		TYR	634	90. 94			1.00 7.5			
ATOM	10888		TYR	634	90. 92			1.00 7.5			
ATOM	10889		TYR	634	88. 649	9 41.660		1.00 8.8			
ATOM	10890		TYR	634	88. 61			1.00 7.8	88 B	C	
ATOM	10891	CZ	TYR	634	89.75			1.00 6.9			
ATOM ATOM	10892 10893	OH C	TYR TYR	634 634	89. 722 88. 967			1.00 8.0			
ATOM	10894	ŏ	TYR	634	88. 038			1.00 13.0 1.00 13.1		C 0	
ATOM	10895	Ň	VAL	635	90.140			1.00 13.1		N	
ATOM	10896	CA	VAL	635	90. 426			1.00 13.3		Č	
ATOM	10897	CB	VAL	635	91.839			1.00 13.2		Č	
ATOM	10898		VAL	635	91, 995			1.00 13.0	6 B	C	
ATOM	10899		VAL	635	92. 894			1.00 8.0		C	
ATOM ATOM	10900	C	VAL	635	89. 412			1.00 13.3		C	
ATOM	10901 10902	O N	VAL THR	635 636	88. 932 89. 091			1.00 15.0		0	
ATOM	10903	CA	THR	636	88. 108			1.00 13.4 1.00 13.7		N	
ATOM	10904	CB	THR	636	87. 788			1.00 15.1		C	
ATOM	10905		THR	636	88. 950			1.00 15.2		Õ	
ATOM	10906		THR	636	86.655			1.00 13.5		Č	
ATOM	10907	C	THR	636	86.792			1.00 14.5	7 B	Č	
ATOM	10908	0	THR	636	86.160			1.00 15.2		0	
ATOM	10909 10910	N CA	SER	637	86.373		41.762	1.00 15.5		N	
ATOM ATOM	10910	CA CB	SER SER	637 637	85.120 84.698			1.00 15.9		C	
ATOM	10912	OG	SER	637	84.303			1.00 16.8 1.00 18.0		C	
ATOM	10913	C	SER	637	85. 195			1.00 16.0		0 C	
ATOM	10914	Ŏ	SER	637	84. 250			1.00 17.8	_	0	
ATOM	10915	N	MET	638	86. 309			1.00 15.6		N	
ATOM	10916	CA	MET	638	86.493		39.052	1.00 15.5		Ċ	
ATOM	10917	CB	MET	638	87.807		38. 272	1.00 15.9		C	
ATOM	10918	CG	MET	638	87. 822		37.067	1.00 17.3		С	• •
ATOM ATOM	10919	SD	MET	638	86. 715		35. 736	1.00 19.14		S	
ATOM	10920 10921	CE C	MET MET	638	87.806		34. 798	1.00 15.23		C	
ATOM	10921		MET	638 638	86. 511 86. 018		40. 093 39. 843	1.00 17.50		C	
ATOM	10923	N	VAL	639	87. 086		41. 260	1.00 17.45 1.00 16.50		O N	
ATOM	10924	ĊA	VAL	639	87.133		42.317	1.00 17.27		C	
ATOM	10925	CB	VAL	639	88.047		43. 480	1.00 16.78		Č	
ATOM	10926	CG1	VAL	639	87.648		44.757	1.00 16.23		Č	

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										(Continued)
					FIC	G. 4-	2 2 4			(0010111404)
ATOM	10927		VAL	639	89. 495 85. 742	35. 335	43. 139	1.00 14.45	В	C
ATOM ATOM	10928 10929	C 0	VAL VAL	639 639	85. 387	34. 919 33. 760	42. 875 43. 081	1.00 17.57 1.00 18.52	B B	C 0
ATOM	10930	N	LEU	640	84. 957	35. 964	43. 124	1.00 16.90	В	N
ATOM	10931	CA	LEU	640	83. 618	35. 766	43.661	1.00 17.42	B	Ĉ
ATOM	10932	CB	LEU	640	82. 978	37.098	44.032	1.00 17.45	В	С
ATOM	10933	CG	LEU	640	83. 512	37.699	45.327	1.00 17.52	В	C .
ATOM	10934		LEU	640	82.743	38.962	45.654	1.00 14.30	В	C
ATOM	10935		LEU	640	83. 378	36.677	46.447	1.00 15.97	В	C
ATOM	10936	Ç.	LEU	640	82. 713	35. 020	42.699	1.00 17.81	В	C
ATOM	10937	0	LEU	640	81.821	34. 284	43.119	1.00 20.73	В	0
ATOM ATOM	10938 10939	N CA	GLY GLY	641 641	82. 952 82. 135	35. 198 34. 526	41.409 40.418	1.00 18.14 1.00 17.61	B B	N
ATOM	10939	CA	GLY	641	82. 758	33. 235	39. 936	1.00 17.01	В	C C
ATOM	10941	0	GLY	641	82. 346	32. 697	38. 911	1.00 17.02	В	0 .
ATOM	10942	Ň	SER	642	83. 735	32. 727	40.683	1.00 17.53	В	Ň
ATOM	10943	CA	SER	642	84. 419	31.497	40. 297	1.00 19.98	B	Ċ
ATOM	10944	CB	SER	642	85.841	31.479	40.864	1.00 20.78	В	C
ATOM	10945	0G	SER	642	85.849	31.088	42.226	1.00 21.56	В	0
ATOM	10946	C	SER	642	83. 691	30. 239	40.755	1.00 21.75	В	C ·
ATOM	10947	0	SER	642	83. 974	29. 147	40. 265	1.00 22.65	<u>B</u> .	0
ATOM	10948	N	GLY	643	82.768	30. 395	41.701	1.00 22.05	В	N
ATOM ATOM	10949 10950	CA	GLY	643	82.023	29. 258	42.210	1.00 22.58	В	C
ATOM	10950	C 0	GLY GLY	643 643	82. 811 82. 460	28. 335 27. 162	43. 130 43. 271	1.00 24.03 1.00 26.05	B B	C
ATOM	10952	N	SER	644	83. 859	28. 849	43. 772	1.00 20.03	В	O. N
ATOM	10953	CA	SER	644	84. 684	28. 024	44. 656	1.00 21.56	В	Č
ATOM	10954	CB	SER	644	86.065	28. 657	44. 833	1.00 21.02	B	č
ATOM	10955	0G	SER	644	85.992	29.798	45.666	1.00 22.35	B	Ö
ATOM	10956	C	SER	644	84.084	27.773	46.037	1.00 21.06	B	Č
ATOM	10957	0	SER	644	84.451	26.807	46.707	1.00 23.51	В	0
ATOM	10958	N	GLY	645	83. 175	28.643	46.469	1.00 19.50	В	N
ATOM	10959	CA	GLY	645	82. 561	28. 485	47. 774	1.00 16.85	В	C
ATOM	10960	C	GLY	645	83.484	28. 868		1.00 18.76	В	C
ATOM ATOM	10961 10962	0 N	GLY	645	83.111	28. 771	50.090	1.00 18.32	В	0
ATOM	10962	N CA	VAL VAL	646 646	84. 691 85. 669	29. 320 29. 695	48. 591 49. 612	1.00 18.97 1.00 18.18	В	N
ATOM	10964	CB	VAL	646	87. 095	29. 718	49.012	1.00 10.10	B B	C C
ATOM	10965		VAL	646	88. 082	30. 202	50.086	1.00 13.30	В	Č
ATOM	10966		VAL	646	87.471	28. 341	48. 516	1.00 17.29	В	č
ATOM	10967	C	VAL	646	85. 433	31.051	50. 266	1.00 18.24	В	č
ATOM	10968	0	VAL	646	85.860	31.270	51.396	1.00 20.76	B	ŏ
ATOM	10969	N	PHE	647	84.763	31.957	49.561	1.00 16.76	B	N
ATOM	10970		PHE	647	84. 525	33. 297	50.082	1.00 16.60	В	C
ATOM	10971		PHE	647	85.066	34. 337	49.094	1.00 16.44	В	C
ATOM	10972		PHE	647	86. 528	34. 204	48. 820	1.00 15.63	В	C
ATOM	10973		PHE	647	87. 455	34. 941	49. 553	1.00 14.72	В	C
ATOM	10974		PHE	647 647	86. 985	33. 320		1.00 14.49	В	C
ATOM	10975	UDI	PHE	041	88. 826	ა4. გიი	49. 317	1.00 16.66	В	С

					FIG. 4-	225			(Cor	itinued)
					1 1 0. 1					
ATOM			2 PHE	647	88. 356 33. 170	47.600	1.00 16.73	В	C	
ATOM		-	PHE	647	89. 278 33. 913	48. 338	1.00 13.35	В	C	
ATOM			PHE	647	83.068 33.604	50. 365	1.00 16.77	В	C	
ATOM			PHE	647	82. 194 33. 328	49. 551	1.00 17.32	В	0	
ATOM	10980	N	LYS	648	82. 819 34. 214	51.515	1.00 16.74	В	N	
ATOM	10981	CA	LYS	648	81. 466 34. 565	51.905	1.00 19.64	В	C	
ATOM ATOM	10982 10983	CB	LYS	648	81.369 34.634	53. 429	1.00 19.84	В	C	
ATOM	10984	CG CD	LYS LYS	648	80.069 35.233	53.911	1.00 21.93	В	C	
ATOM	10985	CE	LYS	648 648	79. 876 35. 060 78. 548 35. 645	55. 393	1.00 23.19	В	C	
ATOM	10986	NZ	LYS	648	78. 548 35. 645 78. 180 35. 150	55. 814 57. 165	1.00 24.97	В	C	
ATOM	10987	C	LYS	648	81. 019 35. 900	51. 308	1.00 31.55 1.00 21.05	. B	N	
ATOM	10988	ŏ	LYS	648	79. 851 36. 070	50. 930	1.00 21.05	B B	C 0	
ATOM	10989	Ň	CYS	649	81. 954 36. 842	51. 237	1.00 20.69	В	N	
ATOM	10990	CA	CYS	649	81. 670 38. 163	50. 711	1.00 21.97	В	Ċ	
ATOM	10991	C	CYS	649	82. 928 38. 811	50. 134	1.00 22.72	B	č	
ATOM	10992	0	CYS	649	84. 054 38. 437	50.477	1.00 23.68	B	ŏ	
ATOM	10993	CB	CYS	649		51.822	1.00 23.52	В	Č	
ATOM	10994	SG	CYS	649		53. 208	1.00 26.89	В	S	•
ATOM	10995	N	GLY	650		49. 267	1.00 20.11	В	N	
ATOM	10996	CA	GLY	650		48.668	1.00 18.42	В	C	
ATOM ATOM	10997 10998	C	GLY	650		48. 308	1.00 18.08	В	C	•
ATOM	10998	O N	GLY ILE	650		48. 135	1.00 18.19	В	0	
ATOM	11000	CA	ILE	651 651		48. 209	1.00 17.42	В	N	
ATOM	11000	CB	ILE	651		47.851	1.00 15.98	В	C	
ATOM	11001		ILE	651		49. 014 48. 589	1.00 15.40 1.00 15.87	B B	C	
ATOM	11003		ILE	651		50. 242	1.00 15.87	В	C	
ATOM	11004	CD1		651		51.411	1.00 13.33	В	C	
ATOM	11005	C	ILE	651		46. 679	1.00 16.40	В	Č	
ATOM	11006	0	ILE	651		46. 754	1.00 16.63	В	ŏ	
ATOM	11007	N	ALA	652		45.608	1.00 16.04	B	Ň	
ATOM	11008	CA	ALA	652	85. 330 45. 4 0 9	44.413	1.00 15.10	B	Ċ	
ATOM	11009	CB	ALA	652		43. 214	1.00 16.38	В	C	
ATOM	11010	C	ALA	652		44. 153	1.00 15.88	В	C	
ATOM	11011	0	ALA	652		43. 895	1.00 14.37	В	0	
ATOM ATOM	11012 11013	N	VAL	653		44. 214	1.00 15.73	В	· C	
ATOM	11013	CA CB	VAL VAL	653		43. 978	1.00 15.50	В	C	
ATOM	11014		VAL	653 653		45.055	1.00 17.97	В	C	
ATOM	11016		VAL	653			1.00 18.06	· B	C	
ATOM	11017	C	VAL	653		46. 446 42. 624	1.00 18.80 1.00 15.11	В	C	
ATOM	11018	ŏ	VAL	653			1.00 15.11	B B	C 0	
ATOM	11019	Ň	ALA	654			1.00 14.41	В	N	
ATOM	11020	CA	ALA	654			1.00 12.10	В	C	
ATOM	11021	CB	ALA	654			1.00 12.15	B	Č	
ATOM	11022	С	ALA	654			1.00 12.48	В	Č	
ATOM	11023	0	ALA	654	88.364 49.500 3		1.00 13.17	B	Ŏ	
ATOM	11024	N	PRO	655			1.00 11.84	В	Ň	
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					FIC	G. 4-	2 2 6			(Continued)
ATOM	11025	CD	PRO	655	85. 273	47.797	40.088	1.00 11.50	В	С
ATOM	11026	CA	PRO	655	87. 247	47.003	38. 954	1.00 11.05	B	č
ATOM	11027	CB	PRO	655	86.399	45.841	39. 436	1.00 11.09	В	č
ATOM	11028	CG	PRO	655	85.030	46. 451	39. 428	1.00 8.50	В	Č
ATOM	11029	C	PRO	655	87. 190	47. 102	37. 447	1.00 10.92	В	č
ATOM	11025	ŏ	PRO	655	86. 383	47.847	36. 896	1.00 10.32	В	ŏ
ATOM	11030	N	VAL	656	88.066	46. 352	36. 791	1.00 9.60	В	N
ATOM	11032	CA	VAL	656	88. 052	46. 250	35. 345	1.00 9.08	В	
ATOM	11032	CB	VAL	656	89. 452	45. 888	34. 790	1.00 7.45	В	C C C
ATOM	11033		VAL	656	89. 336	45. 163	33. 451	1.00 5.90	В	Č
ATOM	11034		VAL	656	90. 249	47. 146	34. 601	1.00 7.63	В	C
ATOM	11036	C	VAL	656	87.107	45.056	35. 224	1.00 10.20	В	C ,
ATOM	11037	Ö	VAL	656	87.157	44. 152	36.058	1.00 10.20	В	0
ATOM	11038	N	SER	657	86. 231	45.038	34. 230	1.00 10.33	В	N
ATOM	11039	CA	SER	657	85. 313	43. 908	34. 115	1.00 11.70	В	L.
ATOM	11033	CB	SER	657	83.867	44. 375	34. 271	1.00 14.03	В	C C
ATOM	11040	OG	SER	657	83.495	45. 242	33. 218	1.00 15.07	В	Õ
ATOM	11041	C	SER	657	85.456	43. 153	32.812	1.00 13.07	В	C
ATOM	11042	ŏ	SER	657	85. 191	41.952	32.743	1.00 14.00	В	Ö
ATOM	11044	N	ARG	658	85.887	43.860	31. 781	1.00 14.15	В	N
ATOM	11044	CA	ARG	658	86.050	43. 277	30. 459	1.00 14.13	В	
ATOM	11046	CB	ARG	658	84. 768	43. 532	29.670	1.00 13.24	В	C
ATOM	11047	CG	ARG	658	84. 763	43. 086	28. 231	1.00 14.22	В	C C C
ATOM	11048	CD	ARG	658	83. 436	43. 470	27. 588	1.00 19.40	В	C
ATOM	11049	NE	ARG	658	83. 475	43. 338	26. 138	1.00 13.40	В	N .
ATOM	11050	CZ	ARG	658	82.868	42.376	25. 454	1.00 23.11	В	C
ATOM	11051	NH1	ARG	658	82.167	41.445	26. 088	1.00 21.95	В	N N
ATOM	11052	NH2	ARG	658	82. 955	42.361	24. 131	1.00 21.33	В	N
ATOM	11053	C	ARG	658	87. 242	44.014	29. 857	1.00 12.76	В	C
ATOM	11054	ŏ	ARG	658	87. 218	45. 239	29. 733	1.00 12.70	В	0
ATOM	11055	Ň	TRP	659	88. 282	43. 283	29.476	1.00 11.05	В	N
ATOM	11056	CA	TRP	659	89. 468	43.942	28. 955	1.00 12.23	В	C
ATOM	11057	CB	TRP	659	90. 578	42.918	28.777	1.00 11.99	В	Č
ATOM	11058	CG	TRP	659	91.026	42. 392	30.112	1.00 13.26	B	C
ATOM	11059		TRP	659	91. 729	43. 120	31. 122	1.00 12.61	В	Č
ATOM	11060		TRP	659	91. 848	42. 271	32. 242	1.00 13.22	В	č
ATOM	11061		TRP	659	92. 268	44. 412	31. 193	1.00 14.19	В	Č
ATOM	11062		TRP	659	90.759	41. 163	30.644	1.00 13.17	В	Č
ATOM	11063		TRP	659	91. 247	41.083	31.920	1.00 13.29	В	Ň
ATOM	11064		TRP	659	92. 489	42.670	33. 424	1.00 13.23	В	C
ATOM	11065		TRP	659	92. 909	44. 810	32. 373	1.00 13.35	В	Č
ATOM	11066		TRP	659	93.011	43. 940	33. 468	1.00 11.92	В	Č
ATOM	11067	C	TRP	659	89. 338	44.840	27. 730	1.00 13.23	В	. Č
ATOM	11068	ŏ	TRP	659	90.118	45. 766	27. 569	1.00 15.25	В	0
ATOM	11069	Ň	GLU	660	88. 361	44. 595	26.871	1.00 13.39	В	N N
ATOM	11070	CA	GLU	660	88. 181	45. 453	25. 708	1.00 14.33	В	C
ATOM	11071	CB	GLU	660	87. 147	44. 854	24. 743	1.00 18.10	В	Č .
ATOM	11072	CG	GLU	660	87. 572	43. 527	24. 130	1.00 21.82	В	Č
ATOM	11073	CD	GLU	660	86. 452	42. 829	23.386	1.00 25.49	В	Č
					00. IUL	10.000	40.000	1.00 40.70	ע	v

					FIC	G. 4	- 227	•		(Contin	nued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11074 11075 11076 11077 11078 11079 11080 11081 11082 11083 11084 11085 11086 11087 11090 11091 11092 11093 11094 11095 11096 11097 11109 11101 11102 11103 11104 11105 11106 11107 11108 11109 11110 11110 11110 11111 11111 11111 11111 11111 11111 1111	OE CON CACE	1 GLUU GLUU GLUU TYRRTYRRTYRRTYRRTYRRTYRRTYRRTYRRTYRRTYR	660 660 660 661 661 661 661 661 661 661	86. 087 85. 929 87. 719 87. 661 87. 371 86. 941 85. 988 84. 599 83. 823 82. 553 84. 061 82. 782 82. 035 88. 146 88. 083 89. 239 90. 411 91. 225 92. 049 93. 379 94. 168 91. 522 92. 297 93. 620 94. 395 91. 095 92. 310 93. 192 93. 961 95. 093 95. 223 95. 869 94. 453 95. 321 95. 464 97. 066 97. 503 98. 865	43. 278 41. 825 46. 833 47. 769 48. 258 48. 119 47. 053 46. 046 49. 045 49. 045 49. 021 48. 699 49. 531 50. 194 51. 030 50. 699 51. 549 49. 615 49. 337 50. 405 51. 741 52. 327 50. 364 49. 658 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371 50. 371	22. 279 23. 914 26. 170 25. 375 27. 450	1.00 29.78 1.00 26.73 1.00 14.88 1.00 14.50 1.00 14.66 1.00 15.13	B B B B B B B B B B B B B B B B B B B	(Continuo o o c o o c o c o c o c o c o c o c o	nued)
ATOM	11119 11120 11121 11122	CG1 CG2 C		665 665 665 665	99. 354 99. 020	51. 263 52. 519 49. 169	27. 206	1. 00 14. 68 1. 00 15. 28 1. 00 15. 25	B B B	C C C	
TIT OM .	11144	U	1 VT	000	99. 972	48. 400	27. 242	1.00 15.22	В	0	

·					FIC	3. 4 <i>-</i>	2 2 8			(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11123 11124 11125 11126 11127 11128 11129 11130 11131 11132 11133 11134 11135 11136 11137 11138 11139 11140	CE1 CD2 CE2 CZ OH C O N CA CB OG1 CG2 C	TYR TYR TYR TYR TYR TYR THR THR THR THR THR THR	666 666 666 666 666 666 666 667 667 667	98. 091 98. 175 97. 504 97. 483 96. 595 96. 583 98. 361 97. 472 97. 471 97. 550 98. 103 96. 401 95. 712 94. 264 93. 617 93. 498 96. 423 96. 713	49. 184 48. 276 48. 896 47. 997 46. 920 46. 089 48. 215 47. 390 46. 332 45. 531 46. 922 45. 895 46. 912 45. 656 44. 624 44. 792 43. 626	28. 154 29. 299 30. 531 31. 751 31. 845 32. 964 32. 809 33. 928 34. 005 35. 131 29. 023 29. 399 28. 365 28. 635 27. 656 28. 635 27. 533 27. 067 27. 323	1. 00 17. 07 1. 00 15. 32 1. 00 13. 28 1. 00 12. 79 1. 00 12. 60 1. 00 12. 83 1. 00 11. 79 1. 00 13. 90 1. 00 15. 26 1. 00 18. 30 1. 00 14. 70 1. 00 13. 70 1. 00 12. 07 1. 00 12. 07 1. 00 10. 21 1. 00 15. 29 1. 00 16. 16	B B B B B B B B B B B B B B B B B B B	N C C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11142 11143 11144 11145 11146 11147 11148 11149 11150 11151 11152 11153 11154 11155 11156 11157 11158	N CA CB CC OE1 OE2 C O N CA CB CC CD NE	GLU GLU GLU GLU GLU GLU GLU ARG ARG ARG ARG ARG ARG	668 668 668 668 668 668 668 669 669 669	96. 707 97. 389 97. 537 96. 231 96. 275 97. 284 95. 284 98. 751 99. 186 99. 418 100. 721 101. 199 102. 498 102. 878 102. 914 102. 549	45. 372 44. 672 45. 612 45. 808 46. 928 47. 054 47. 679 44. 127 43. 079 44. 827 44. 392 45. 291 44. 828 45. 766 47. 149 48. 196	25. 906 24. 823 23. 625 22. 867 21. 850 21. 123 21. 767 25. 247 24. 766 26. 158 26. 640 27. 785 28. 451 29. 583 29. 122 29. 856	1.00 16.99 1.00 17.50 1.00 21.31 1.00 22.06 1.00 25.39 1.00 22.03 1.00 17.77 1.00 19.28 1.00 17.62 1.00 17.00 1.00 17.11 1.00 15.99 1.00 15.35 1.00 16.25 1.00 16.96	B B B B B B B B B B B B B B B B B B B	N C C C O N C C C N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11158 11159 11160 11161 11162 11163 11164 11165 11166 11167 11168 11169 11170 11171	NH2 C O N CA CB CG CD1 CE1 CD2 CE2	ARG ARG TYR TYR TYR TYR TYR TYR TYR TYR TYR	669 669 669 670 670 670 670 670 670 670	102. 115 102. 602 100. 633 101. 523 99. 539 99. 357 98. 823 99. 571 98. 978 99. 680 100. 894 101. 608 100. 998 101. 713	48. 023 49. 417 42. 960 42. 141 42. 655 41. 333 41. 465 42. 491 43. 706 44. 676 42. 268 43. 232 44. 433 45. 403	31. 101 29. 340 27. 140 26. 899 27. 825 28. 385 29. 810 30. 631 30. 973 31. 676 31. 024 31. 732 32. 051 32. 714	1. 00 16. 86 1. 00 14. 86 1. 00 17. 70 1. 00 17. 72 1. 00 17. 60 1. 00 16. 56 1. 00 15. 82 1. 00 15. 47 1. 00 14. 36 1. 00 15. 93 1. 00 15. 78 1. 00 15. 30 1. 00 15. 22	B B B B B B B B B	N C O N C C C C C C C

					FIG. 4-229	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11172 11173 11174 11175 11176 11177 11178 11179 11180 11181 11182 11183 11184 11185 11186 11187 11188 11190 11191 11192 11193 11194 11195 11196 11197	CD2 C O N CD CA CB	TYR TYR MET MET MET MET MET MET MET MET MET LEU LEU LEU LEU LEU PRO PRO PRO PRO	670 670 671 671 671 671 671 671 672 672 672 673 673 673 673 673 673 674 674	FIG. 4 - 229 98. 435	C O N C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11198 11199 11200 11201 11202 11203 11204 11205 11206 11207 11208 11209 11210 11211 11212 11213 11214 11215 11216 11217 11218	CG C O N CA CB OG1 CG2 C O N CD CA CB CG C O N CA CB CG	PRO PRO PRO THR THR THR THR THR PRO PRO PRO PRO GLU GLU GLU GLU GLU	674 674 675 675 675 675 675 676 676 676 676 677 677	90. 845 43. 664 20. 941 1. 00 18. 24 B 90. 589 41. 155 19. 183 1. 00 21. 53 B 89. 470 41. 561 18. 884 1. 00 20. 30 B 91. 378 40. 505 18. 335 1. 00 23. 61 B 90. 973 40. 176 16. 975 1. 00 23. 43 B 92. 045 40. 560 15. 957 1. 00 22. 99 B 93. 221 39. 783 16. 200 1. 00 24. 15 B 92. 386 42. 039 16. 062 1. 00 21. 26 B 90. 825 38. 668 16. 931 1. 00 25. 46 B 91. 424 37. 952 17. 736 1. 00 25. 82 B 90. 023 38. 160 15. 991 1. 00 26. 60 B 89. 130 38. 885 15. 074 1. 00 25. 76 B 89. 823 36. 714 15. 877 1. 00 26. 64 B 88. 860 36. 599 14. 702 1. 00 25. 84 B 88. 066 37. 859 14. 801 1. 00 24. 99 B 91. 135 <	C C C C C C C C C C C C C C C C C C C
ATOM	11218 11219 11220	CD OE1	GLU	677 677	94. 177 35. 077 12. 294 1. 00 41. 76 B 92. 984 34. 204 11. 897 1. 00 46. 15 B 92. 234 34. 610 10. 980 1. 00 49. 52 B	C

					T I /	G. 4-	230			(Continued)
					r I (J. 4 -	230			
ATOM	11221		GLU	677	92. 789	33. 121	12.503	1.00 46.47	В	0 .
ATOM	11222	C	GLU	677	94. 382	36.174	15. 563	1.00 31.51	В	- C
ATOM	11223	0	GLU	677	95.565	35. 938	15.305	1.00 31.18	В	0
ATOM	11224	N	ASP	678	94.003	36.680	16.730	1.00 29.04	В	N
ATOM	11225	CA	ASP	678	95.005	36.896	17.756	1.00 26.71	В	C
ATOM	11226	CB	ASP	678	95. 359	38. 374	17. 917	1.00 25.30	В	C
ATOM	11227	CG	ASP	678	96.500	38. 586	18. 902	1.00 26.53	В	C
ATOM	11228		ASP	678	97.004	39. 721	19.008	1.00 29.18	В	0
ATOM	11229		ASP	678	96.900	37.612	19.579	1.00 24.47 1.00 25.24	В	0
ATOM ATOM	11230	C	ASP	678	94. 586	36. 325	19.098		В	C
ATOM	11231 11232	0 N	ASP ASN	678 679	94. 946 93. 814	35. 200 37. 082	19. 426 19. 871	1.00 26.23 1.00 24.14	В	0 N
ATOM	11232	CA	ASN.	679	93. 418	36.608	21. 186	1.00 24.14	B B	N C
ATOM	11234	CB	ASN.	679	94. 456	37. 089	22. 217	1.00 23.05	В	C C
ATOM	11235	CG	ASN	679	94. 390	36. 323	23. 524	1.00 23.03	В	Č
ATOM	11236		ASN	679	94. 644	36. 880	24. 592	1.00 21.44	В	0
ATOM	11237		ASN	679	94. 059	35. 037	23. 448	1.00 21.44	В	N N
ATOM	11238	C	ASN	679	92. 019	37. 061	21. 596	1.00 21.85	В	Ċ
ATOM	11239	ŏ	ASN	679	91. 727	37. 174	22. 785	1.00 21.56	В	Ö
ATOM	11240	Ň	LEU	680	91. 153	37. 316	20.619	1.00 22.96	B	N
ATOM	11241	ĊA	LEU	680	89. 783	37. 750	20. 913	1.00 22.05	В	Ċ
ATOM	11242	CB	LEU	680	88. 999	37. 967	19.617	1.00 20.94	B	č
ATOM	11243	CG	LEU	680	87.524	38. 379	19.734	1.00 20.98	B	č
ATOM	11244		LEU	680	87. 385	39.671	20.539	1.00 21.18	B	Č
ATOM	11245	CD2	LEU	680	86.946	38.567	18.348	1.00 17.15	В	Ċ
ATOM	11246	C	LEU	680	89.031	36.762	21.805	1.00 22.36	В	C
ATOM	11247	0	LEU	680	88.316	37.171	22.718	1.00 23.81	В	0
ATOM	11248	N	ASP	681	89. 193	35.466	21.555	1.00 22.95	В	N
ATOM	11249	CA	ASP	681	88. 502	34.469	22.371	1.00 24.27	В	· C
ATOM	11250	CB	ASP	681	88.910	33.048	21.980	1.00 24.73	В	C
ATOM	11251	CG	ASP	681	88. 270	32.587	20.695	1.00 25.98	В	C
ATOM	11252		ASP	681	87. 453	33. 334	20.116	1.00 28.21	В	0
ATOM	11253		ASP	681	88. 587	31.462	20. 259	1.00 28.60	В	0
ATOM	11254	C	ASP	681	88. 754	34.655	23.862	1.00 23.99	В	C
ATOM	11255	0	ASP	681	87. 816	34.640	24.660	1.00 24.77	В	0
ATOM	11256	N	HIS	682	90.014	34. 819	24. 252	1.00 22.66	В	N
ATOM	11257	CA	HIS	682	90. 289	34. 998	25.667	1.00 22.62	В	C
ATOM	11258	CB	HIS	682	91.775	34.867	25. 981	1.00 23.03	В	C
ATOM	11259	CG	HIS	682	92.063	34. 898	27. 448	1.00 25.79	В	C
ATOM	11260		HIS	682	92. 844	35. 718	28. 190	1.00 26.73	В	C
ATOM	11261		HIS	682	91.458	34. 035	28. 338	1.00 25.30	В	N
ATOM	11262		HIS	682	91.852	34. 326	29.565	1.00 26.50	В	C
ATOM	11263		HIS	682	92.693	35. 344	29.504	1.00 26.09	В	N
ATOM	11264	C	HIS	682	89.775	36. 344	26. 175	1.00 21.71	В	C
ATOM	11265	0 N	HIS	682	89.412	36. 465	27. 345	1.00 20.98	В	0
ATOM	11266	N	TYR	683	89. 753	37. 355	25. 307	1.00 19.91	В	N
ATOM ATOM	11267 11268	CA CB	TYR TYR	683 683	89. 232	38.657	25.707		В	C
ATOM	11269	CG	TYR	683	89. 226	39.646	24. 542	1.00 16.55	В	C
UIOM	11602	VU	111/	UQJ	90.419	40. 574	24.472	1.00 16.85	В	C

					FIG. 4-232		(Continued)
ATOM ATOM ATOM ATOM ATOM	11319 11320 11321 11322 11323	CB CG SD CE C	MET MET MET MET MET	689 689 689 689	79.519 40.287 32.010 1.00 14.10 79.359 41.793 32.217 1.00 18.18 80.817 42.684 32.849 1.00 21.67 81.693 43.067 31.308 1.00 19.11 79.429 38.040 33.080 1.00 13.66	B B B B	C C S C
ATOM ATOM ATOM ATOM ATOM ATOM	11324 11325 11326 11327 11328 11329	O N CA CB OG	MET SER SER SER SER	689 690 690 690	78. 398 37. 597 33. 586 1. 00 14. 01 80. 246 37. 290 32. 356 1. 00 14. 32 79. 939 35. 887 32. 087 1. 00 16. 68 81. 018 35. 259 31. 199 1. 00 18. 28 82. 225 35. 062 31. 923 1. 00 23. 11	B B B B	O N C C O
ATOM ATOM ATOM ATOM ATOM	11330 11331 11332 11333 11334	C O N CA CB CG	SER SER ARG ARG ARG ARG	690 690 691 691 691	79. 771 35. 019 33. 328 1. 00 15. 55 79. 212 33. 927 33. 234 1. 00 16. 21 80. 238 35. 502 34. 478 1. 00 14. 35 80. 155 34. 741 35. 727 1. 00 15. 38 81. 491 34. 821 36. 478 1. 00 16. 76 82. 697 34. 414 35. 652 1. 00 19. 96	B B B B B	C O N C C C
ATOM ATOM ATOM ATOM ATOM	11335 11336 11337 11338 11339	CD NE CZ NH1 NH2	ARG ARG ARG ARG ARG	691 691 691 691	83. 972 34. 339 36. 483 1. 00 21. 36 85. 061 33. 725 35. 726 1. 00 23. 56 86. 196 33. 274 36. 256 1. 00 26. 24 86. 418 33. 358 37. 567 1. 00 23. 55 87. 114 32. 728 35. 468 1. 00 26. 33	B B B B	C N C N N
ATOM ATOM ATOM ATOM ATOM ATOM	11340 11341 11342 11343 11344 11345	C O N CA CB C	ARG ARG ALA ALA ALA ALA	691 691 692 692 692 692	79. 049 35. 187 36. 679 1. 00 15. 48 78. 986 34. 713 37. 817 1. 00 14. 38 78. 178 36. 081 36. 220 1. 00 14. 78 77. 111 36. 618 37. 064 1. 00 16. 42 76. 105 37. 383 36. 198 1. 00 16. 75	B B B B	C . O . N . C . C . C
ATOM ATOM ATOM ATOM ATOM	11346 11347 11348 11349 11350	O N CA CB	ALA GLU GLU GLU GLU	692 693 693 693 693	76. 375 35. 624 37. 977 1. 00 17. 17 76. 331 35. 814 39. 191 1. 00 16. 75 75. 803 34. 571 37. 404 1. 00 19. 44 75. 062 33. 589 38. 191 1. 00 22. 16 74. 570 32. 443 37. 299 1. 00 26. 71 73. 251 32. 745 36. 598 1. 00 33. 79	B B B B	C O N C C C
ATOM ATOM ATOM ATOM ATOM	11351 11352 11353 11354 11355	CD OE1	GLU GLU GLU GLU GLU	693 693 693 693	73. 017 31. 873 35. 379 1. 00 38. 47 72. 984 30. 632 35. 531 1. 00 40. 41 72. 870 32. 433 34. 266 1. 00 41. 15 75. 827 33. 022 39. 369 1. 00 22. 08 75. 244 32. 761 40. 418 1. 00 24. 44	B B B B	C 0 0 C
ATOM ATOM ATOM ATOM	11356 11357 11358 11359 11360	N CA CB CG OD1	ASN ASN ASN ASN	694 694 694 694	77. 127 32. 824 39. 215 1. 00 21. 66 77. 907 32. 282 40. 320 1. 00 22. 61 79. 324 31. 924 39. 861 1. 00 20. 93 79. 359 30. 654 39. 048 1. 00 19. 32 80. 284 30. 420 38. 278 1. 00 19. 68	B B B B	N C C C
ATOM ATOM ATOM ATOM ATOM ATOM	11361 11362 11363 11364 11365 11366	ND2 C O N CA CB	ASN ASN PHE PHE PHE	694 694 694 695 695	78. 348 29. 818 39. 224 1. 00 18. 34 77. 975 33. 234 41. 500 1. 00 22. 99 78. 650 32. 946 42. 479 1. 00 25. 59 77. 283 34. 366 41. 419 1. 00 22. 83 77. 299 35. 316 42. 531 1. 00 23. 74 77. 205 36. 772 42. 041 1. 00 20. 88	B B B B	N C O N C
ATOM	11367	CG	PHE	695	77. 205 36. 772 42. 041 1. 00 20. 88 78. 533 37. 397 41. 695 1. 00 19. 06	B B	C C

					Tr 1 C 4 . 9 9 9	(Continued)
					FIG. 4-233	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11368 11369 11370 11371 11372 11373 11374	CD2 CE1	PHE PHE PHE PHE PHE PHE PHE	695 695 695 695 695	79. 211 37. 042 40. 533 1. 00 19. 50 B 79. 096 38. 365 42. 523 1. 00 19. 69 B 80. 431 37. 647 40. 200 1. 00 18. 29 B 80. 316 38. 977 42. 199 1. 00 18. 53 B 80. 982 38. 615 41. 033 1. 00 17. 35 B 76. 146 35. 052 43. 483 1. 00 24. 37 B	C C C C
ATOM ATOM ATOM ATOM	11375 11376 11377 11378	N CA CB CG	LYS LYS LYS LYS	695 696 696 696	76. 090 35. 636 44. 566 1. 00 25. 67 B 75. 230 34. 173 43. 089 1. 00 24. 40 B 74. 074 33. 880 43. 926 1. 00 25. 82 B 73. 173 32. 813 43. 280 1. 00 27. 75 B 72. 076 32. 281 44. 228 1. 00 30. 02 B	O N C C C
ATOM ATOM ATOM ATOM	11379 11380 11381 11382	CD CE NZ C	LYS LYS LYS LYS	696 696 696 696	70. 680 32. 287 43. 615 1. 00 31. 63 B 70. 137 33. 705 43. 421 1. 00 35. 45 B 69. 903 34. 438 44. 705 1. 00 35. 47 B 74. 402 33. 459 45. 348 1. 00 24. 85 B	C C N C
ATOM ATOM ATOM ATOM ATOM	11383 11384 11385 11386 11387	O N CA CB CG	LYS GLN GLN GLN GLN	696 697 697 697	73. 583 33. 641 46. 242 1. 00 24. 94 B 75. 587 32. 907 45. 577 1. 00 25. 99 B 75. 920 32. 481 46. 931 1. 00 27. 33 B 76. 355 31. 010 46. 941 1. 00 29. 90 B 75. 290 30. 025 46. 444 1. 00 30. 66 B	O N C C C C
ATOM ATOM ATOM ATOM	11388 11389 11390 11391	CD OE1 NE2 C	GLN GLN GLN GLN	697 697 697 697	75. 565 28. 593 46. 889 1. 00 30. 92 B 75. 381 28. 245 48. 065 1. 00 31. 54 B 76. 019 27. 761 45. 958 1. 00 26. 21 B 76. 964 33. 322 47. 662 1. 00 26. 04 B	C O N C
ATOM ATOM ATOM ATOM ATOM	11392 11393 11394 11395 11396	O N CA CB CG1	GLN VAL VAL VAL VAL	697 698 698 698	77. 620 32. 833 48. 580 1. 00 28. 31 B 77. 125 34. 580 47. 270 1. 00 23. 16 B 78. 085 35. 445 47. 947 1. 00 21. 23 B 79. 411 35. 596 47. 156 1. 00 20. 63 B 80. 033 34. 238 46. 901 1. 00 17. 19 B	O N C C
ATOM ATOM ATOM ATOM	11397 11398 11399 11400	CG2 C O N	VAL VAL VAL GLU	698 698 698 699	79. 161 36. 335 45. 853 1. 00 18. 36 B 77. 496 36. 829 48. 118 1. 00 21. 50 B 76. 571 37. 207 47. 404 1. 00 23. 06 B 78. 018 37. 579 49. 078 1. 00 21. 31 B	C C C O N
ATOM ATOM ATOM ATOM ATOM	11401 11402 11403 11404 11405	CA CB CG CD OE1	GLU GLU GLU GLU GLU	699 699 699 699	77. 563 38. 945 49. 290 1. 00 21. 42 B 77. 465 39. 246 50. 785 1. 00 22. 73 B 76. 396 38. 403 51. 461 1. 00 26. 07 B 76. 547 38. 346 52. 961 1. 00 29. 09 B	C C C
ATOM ATOM ATOM ATOM	11406 11407 11408 11409		GLU GLU GLU TYR	699 699 699 700	76. 343 39. 387 53. 624 1. 00 31. 29 B 76. 876 37. 254 53. 476 1. 00 31. 07 B 78. 610 39. 810 48. 593 1. 00 21. 23 B 79. 802 39. 751 48. 905 1. 00 21. 45 B 78. 148 40. 594 47. 630 1. 00 19. 47 B	0 0 C 0 N
ATOM ATOM ATOM ATOM	11410 11411 11412 11413	CA CB CG CD1	TYR TYR TYR TYR	700 700 700 700	79. 012 41. 428 46. 818 1. 00 18. 26 B 78. 830 41. 001 45. 368 1. 00 18. 24 B 79. 678 41. 685 44. 330 1. 00 18. 56 B 81. 071 41. 698 44. 422 1. 00 17. 75 B	C C C
ATOM ATOM ATOM	11414 11415 11416	CE1 CD2 CE2	TYR	700 700 700	81. 856 42. 206 43. 378 1. 00 17. 99 B 79. 088 42. 209 43. 181 1. 00 19. 07 79. 852 42. 715 42. 143 1. 00 19. 54 B	C C C

										(Continued)
					FIC	G. 4-	234			(Continued)
					`			**		
ATOM	11417	CZ	TYR	700	81.231	42. 707	42.241	1.00 19.61	В	C
ATOM	11418	0H	TYR	700	81.964	43. 170	41.176	1.00 20.17	В	0
ATOM	11419	C	TYR	700	78.697	42.902	46.972	1.00 18.20	В	C
ATOM	11420	0	TYR	700	77. 534	43. 288	47.006	1.00 19.67	В	0
ATOM	11421	N	LEU	701	79. 748	43.714	47.078	1.00 16.71	В	N
ATOM	11422	CA	LEU	701	79.628	45. 157	47.198	1.00 15.24	В	C
ATOM	11423	CB	LEU	701	80. 102	45.624	48.573	1.00 14.82	В	C
ATOM	11424	CG	LEU	701	80. 195	47. 141	48.768	1.00 15.42	В	C
ATOM	11425		LEU	701	78. 926	47. 810	48. 280	1.00 16.37	В	C
ATOM	11426		LEU	701	80.449	47. 456	50. 233	1.00 13.32	В	C
ATOM ATOM	11427	C	LEU	701	80. 491	45. 770	46.095	1.00 16.15	В	C
ATOM	11428 11429	0 N	LEU LEU	701 702	81. 714 79. 829		46. 082 45. 167	1.00 16.12	В	0
ATOM	11425	CA	LEU	702 702	80.467	46. 450 47. 073	43.107	1.00 14.91 1.00 13.94	B B	N C
ATOM	11431	CB	LEU	702	79. 730	46. 627	42. 753	1.00 15.12	В	Č
ATOM	11432	CG	LEU	702	80.119	47. 175	41. 383	1.00 15.12	В	Č
ATOM	11433		LEU	702	81.555	46. 814	41.050	1.00 14.64	В	č
ATOM	11434		LEU	702	79. 173	46. 593	40. 354	1.00 16.45	В	č
ATOM	11435	C	LEU	702	80.419	48. 590	44. 169	1.00 14.21	B	č
ATOM	11436	0	LEU	702	79.346	49. 166	44. 314	1.00 14.96	B	Ö
ATOM	11437	N	ILE	703	81.591	49. 220	44.132	1.00 13.90	B	N
ATOM	11438	CA	ILE	703	81.737	50.662	44. 294	1.00 13.91	В	C
ATOM	11439	CB	ILE	703	82.543	50.967	45.578	1.00 13.87	В	C
ATOM	11440		ILE	703	82.693	52.491	45.775	1.00 15.37	В	С
ATOM	11441		ILE	703	81.869	50. 308	46.782	1.00 12.11	В	C
ATOM	11442	CD1	ILE	703	82. 714	50. 328	48.047	1.00 7.95	В	C
ATOM	11443	C	ILE	703	82.495	51. 251	43. 101	1.00 15.43	В	C
ATOM	11444	0	ILE	703	83. 379	50.600	42.548	1.00 17.12	В	0
ATOM	11445	N	HIS	704	82. 175	52. 484	42.714	1.00 14.44	В	N
ATOM ATOM	11446 11447	CA	HIS	704	82.866	53.098	41.579	1.00 14.11	В	C
ATOM	11448	CB CG	HIS HIS	704	82. 483	52. 356 52. 386	40. 288	1.00 12.85	В	C
ATOM	11449		HIS	704 704	83. 539 84. 363	52. 360 53. 377	39. 224 38. 806	1.00 13.44	В	C
ATOM	11445		HIS	704	83. 827	51. 293	38. 435	1.00 12.54 1.00 12.00	B B	C N
ATOM	11451		HIS	704	84. 782	51.607	37. 578	1.00 12.00	В	C
ATOM	11452		HIS	704	85. 125	52. 865	37. 782	1.00 10.03	В	N N
ATOM	11453	C	HIS	704	82. 533	54. 584	41.457	1.00 12.00	В	Č
ATOM	11454	Ŏ	HIS	704	81.420	55.007	41.770	1.00 15.67	B	<u> </u>
ATOM	11455	N	GLY	705	83. 513	55. 372	41.027	1.00 10.99	B	Ň
ATOM	11456	CA	GLY	705	83. 308	56. 798	40.860	1.00 10.39	B	Ĉ
ATOM	11457	C	GLY	705	82.807		39.457	1.00 10.13	B	č
ATOM	11458	0	GLY	705	83. 326	56.536	38.483	1.00 11.85	B	Ö
ATOM	11459	N	THR	706	81.805	57.942	39.347	1.00 10.36	В	N
ATOM	11460	CA	THR	706	81. 215	58. 272	38.054	1.00 9.96	В	C
ATOM	11461	CB	THR	706	79. 935	59.072	38. 232	1.00 6.56	В	C
ATOM	11462	0G1	THR	706	80. 251	60.367	38.739	1.00 8.64	В	0
ATOM	11463	CG2	THR	706	79. 025	58. 372	39. 215	1.00 8.26	В	Č .
ATOM	11464	C	THR	706	82. 145	59.052	37. 147	1.00 11.88	В	C
ATOM	11465	0	THR	706	81.994	59.018	35.927	1.00 13.83	В	0

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					FIG. 4-235	(Continued)
ATOM	11466	N	ALA	707	83. 114 59. 741 37. 739 1. 00 13. 21 B	N
ATOM	11467	CA	ALA	707	84. 075 60. 522 36. 969 1. 00 14. 57 B	C
ATOM	11468	CB	ALA	707	84. 277 61. 881 37. 626 1. 00 17. 64 B	C
ATOM	11469	C	ALA	707	85. 427 59. 823 36. 802 1. 00 13. 77 B 86. 445 60. 484 36. 639 1. 00 14. 15 B	C
ATOM	11470	0	ALA	707		O
ATOM	11471	N	ASP	708	85. 435 58. 494 36. 839 1. 00 13. 35 B	N
ATOM	11472	CA	ASP	708	86. 667 57. 721 36. 685 1. 00 12. 65 B	
ATOM ATOM	11473 11474	CB CG	ASP ASP	708 708	86. 439 56. 285 37. 188 1. 00 12. 24 B 87. 737 55. 536 37. 453 1. 00 10. 05 B	C C C
ATOM ATOM	11475 11476	0D1	ASP ASP	708 708	88. 738 55. 775 36. 749 1. 00 11. 19 B 87. 751 54. 686 38. 362 1. 00 9. 31 B	0
ATOM	11477	C	ASP	708	87. 091 57. 696 35. 202 1. 00 13. 18 B	0
ATOM	11478	0	ASP	708		C
ATOM	11479	N	ASP	709	88. 156 58. 423 34. 891 1. 00 12. 80 B	0
ATOM	11480	CA	ASP	709		N
ATOM	11481	CB	ASP	709	89. 442 59. 825 33. 397 1. 00 11. 74 B	C
ATOM	11482	CG	ASP	709		C
ATOM ATOM	11483 11484	0D1	ASP ASP	709 709	90. 612 59. 912 34. 366 1. 00 9. 63 B 91. 704 59. 385 34. 058 1. 00 2. 39 B 90. 419 60. 499 35. 451 1. 00 11. 84 B	C 0
ATOM ATOM	11485 11486	C 0	ASP ASP	709 709	89. 605 57. 366 33. 167 1. 00 14. 57 89. 896 57. 136 31. 987 1. 00 16. 47	0 C 0
ATOM	11487	N	ASN	710	90.076 56.652 34.182 1.00 13.58 B 90.981 55.524 33.990 1.00 13.56 B	N
ATOM	11488	CA	ASN	710		C
ATOM	11489	CB	ASN	710	91. 841 55. 385 35. 243 1. 00 13. 26 B 92. 987 54. 440 35. 059 1. 00 12. 07 B	C
ATOM	11490	CG	ASN	710		C
ATOM ATOM	11491 11492	ND2	ASN ASN	710 710	93. 951 54. 478 35. 821 1. 00 16. 69 B 92. 898 53. 578 34. 058 1. 00 8. 28 B	O N
ATOM	11493	0	ASN	710	90. 177 54. 236 33. 724 1. 00 14. 26 B 90. 142 53. 737 32. 598 1. 00 14. 29 B	C
ATOM	11494	C	ASN	710		0
ATOM	11495	N	VAL	711	89. 560 53. 692 34. 773 1. 00 13. 24 B	N
ATOM	11496	CA	VAL	711	88. 715 52. 511 34. 652 1. 00 12. 56 B	C
ATOM	11497		VAL	711	88. 835 51. 585 35. 868 1. 00 11. 72 B	C
ATOM	11498		VAL	711	88. 048 50. 311 35. 624 1. 00 7. 36 B	C
ATOM ATOM	11499 11500	C	VAL VAL	711 711	90. 287 51. 274 36. 141 1. 00 13. 94 B 87. 315 53. 119 34. 645 1. 00 14. 01 B	C . C
ATOM	11501	O	VAL	711	86. 768 53. 471 35. 694 1. 00 13. 52 B 86. 746 53. 249 33. 456 1. 00 13. 66 B	O
ATOM	11502	N	HIS	712		N
ATOM ATOM ATOM	11503 11504 11505	CA CB CG	HIS HIS HIS	712 712	85. 440 53. 869 33. 290 1. 00 13. 44 B 85. 132 53. 956 31. 794 1. 00 12. 94 B	C
ATOM ATOM	11505 11506 11507	CD2	HIS HIS	712 712 712	86. 219 54. 613 31. 001 1. 00 14. 38 B 87. 137 55. 549 31. 352 1. 00 15. 50 B 86. 477 54. 299 29. 684 1. 00 15. 76 B	C C
ATOM ATOM	11508 11509	CE1	HIS HIS	712 712 712	87. 510 55. 009 29. 258 1. 00 17. 42 B	N C
ATOM ATOM	11510 11511	C	HIS HIS	712 712	84. 293 53. 205 34. 048 1. 00 13. 09 B	N C ·
ATOM	11512	N	PHE	713	83. 420 54. 041 34. 594 1. 00 13. 27 B	0
ATOM	11513	CA	PHE	713		N
ATOM	11514	CB	PHE	713	82. 253 53. 586 35. 335 1. 00 15. 36 B 81. 288 54. 759 35. 530 1. 00 15. 17 B	C C

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					r ı	G.	4 -	2 3 6					
ATOM	11515	CG	PHE	713	80. 15		. 464	36. 461		16.61	В	C	
ATOM	11516		PHE	713	80.34		. 508	37. 841		14.51	В	C	
ATOM ATOM	11517 11518		PHE	713 713	78. 90 79. 30		. 111	35. 962 38. 710		15. 42 14. 71	B B	C C	
ATOM	11516		PHE PHE	713	79. 30 77. 84		8.803	36. 829		15. 24	В	C	
ATOM	11519	CZ	PHE	713	78. 05		. 849	38. 204		13. 41	В	Č	
ATOM	11520	C	PHE	713	81. 58		. 486	34. 499		16. 62	В	č	
ATOM	11522	ŏ	PHE	713	81.01		. 527	35. 031		16. 48	В	ŏ	
ATOM	11523	Ň	GLN	714	81.67		. 649	33. 181		15. 73	B	Ň	
ATOM	11524	CA	GLN	714	81.12		. 699	32. 228		16.08	В	C	
ATOM	11525	CB	GLN	714	81.75		. 923	30.857	1.00	14.90	В	C	
ATOM	11526	CG	GLN	714	81.69		. 703	29.946		16.13	В	C	
ATOM	11527	CD	GLN	714	82.66		. 811	28.770		15.37	В	С	
ATOM	11528		GLN	714	83. 82		.167	28. 943		15.11	В	0	
ATOM	11529		GLN	714	82. 18		. 493	27.577		15.35	В	N	
ATOM	11530	C	GLN	714	81.37		. 256	32.650		16. 29	В	C	
ATOM ATOM	11531 11532	O N	GLN GLN	714	80. 51		. 389	32. 487		17.82	В	0 N	
ATOM ATOM	11532	CA	GLN	715 715	82. 55 82. 90		. 997	33. 192 33. 593		14. 60 14. 55	B B	N C	
ATOM	11534	CB	GLN	715	84. 39		581	33. 926		16. 22	В	C	
ATOM	11535	CG	GLN	715	85. 27		. 086	32.767		16. 01	В	č	
ATOM	11536	CD	GLN	715	86. 50		. 247	32. 537		14. 28	В	č	
ATOM	11537		GLN	715	86. 47		. 029	32.674		17.54	B	Ŏ	
ATOM	11538		GLN	715	87.60		. 889	32. 155		12.78	В	N	
ATOM	11539	C	GLN	715	82.03	1 48	. 134	34.746	1.00	14.99	В	C	
ATOM	11540	0	GLN	715	81.61		.967	34.749		13.70	В	0	
ATOM	11541	N	SER	716	81.74		. 002	35. 714		12.14	В	N	
ATOM	11542	CA	SER	716	80. 89		. 602	36. 829		11.18	В	C	
ATOM	11543	CB	SER	716	81.05		. 544	38. 028		11.19	В	C	•
ATOM ATOM	11544 11545	OG C	SER SER	716 716	82. 27		. 295	38. 700		13.48	В	0	
ATOM	11546	0	SER	716	79. 43 78. 68		. 570 . 692	36. 394 36. 814	1.00 1.00	9. 18 5. 81	B B	C 0	
ATOM	11547	N	ALA	717	79. 02		. 517	35. 552	1.00	8. 69	В	N N	
ATOM	11548	CA	ALA	717	77. 63		. 537	35. 083		10.91	В	C	
ATOM	11549	CB	ALA	717	77. 40		. 708	34. 143		10.07	В	Č	
ATOM	11550	C	ALA	717	77. 30		. 219	34. 382		10.72	B	Č	
ATOM	11551	0	ALA	717	76. 21		. 696	34.539		14.08	В	Ō	
ATOM	11552	N	GLN	718	78. 25	2 47	. 682	33.623	1.00	10.89	В	N	
ATOM	11553	CA	GLN	718	78.05		. 417	32. 928	1.00	10.32	В	С	
ATOM	11554	CB	GLN	718	79. 13		. 224	31.858	1.00	8. 83	В	C	
ATOM	11555	CG	GLN	718	79.07		. 232	30. 722	1.00	6.53	В	C	
ATOM	11556	CD	GLN	718	78.00		. 900	29. 691	1.00	8. 70	В	C	
ATOM	11557		GLN	718	76. 97		. 319	30. 012		13. 43	В	0	
ATOM ATOM	11558 11559		GLN GLN	718 718	78. 24		. 278	28. 449		11.12	В	N	
ATOM	11560	C 0	GLN	718	78. 05 77. 35		. 235 . 248	33. 908 33. 695		10.68 13.48	B	C	
ATOM	11561	·N	ILE	719	78. 83		. 248 . 320	33. 695 34. 981		12. 24	B B	O N	
ATOM	11562	CA	ILE	719	78. 85		. 226	35. 953		12. 41	В	C	
ATOM	11563	CB	ILE	719	79.89		. 434			12. 88	В	Č	
VIII		0.0					- 101	01.010		00	b	v	

						_	0.000			(Continued)
					FIC	G. 4 -	2 3.8		•	
ATOM ATOM	11613 11614	CA CB	VAL VAL	726 726	70. 409 71. 727	36. 726 35. 920	36. 329 36. 392	1.00 17.93 1.00 19.28	B B	C C
ATOM ATOM	11615 11616		VAL VAL	726 726	72. 246 72. 763	35.672 36.660	34. 994 37. 238	1.00 19.33 1.00 19.80	B B	C C
ATOM	11617	C	VAL	726	69. 789	36.741	37. 723	1.00 17.35	В	C
ATOM ATOM	11618 11619	O N	VAL GLY	726 727	69. 858 69. 198	35. 756 37. 875	38. 463 38. 081	1.00 16.63 1.00 17.14	B B	O N
ATOM	11620	CA	GLY	727	68.548	38.012	39.370	1.00 15.42	В	C
ATOM	11621	C	GLY	727	69.387	37.856	40.626 41.559	1.00 15.90 1.00 17.97	B B	C .
ATOM ATOM	11622 11623	O N	GLY VAL	727 728	68. 961 70. 568	37. 182 38. 462	41. 559	1.00 17.97	В	O N
ATOM	11624	CA	VAL	728	71.389	38.357	41.876	1.00 14.10	В	C .
ATOM	11625	CB	VAL	728	72.859	37. 972	41.574	1.00 14.97	В	C
ATOM ATOM	11626 11627		VAL VAL	728 728	73. 693 72. 954	38. 145 36. 514	42.829 41.109	1.00 13.51 1.00 15.40	B B	C C
ATOM	11628	C	VAL	728	71.396	39.687	42.603	1.00 14.73	B	č
ATOM	11629	0	VAL	728	71. 738	40.714	42.025	1.00 14.56	В	0
ATOM ATOM	11630 11631	N CA	ASP ASP	729 729	71.007 70.998	39. 672 40. 896	43.872 44.646	1.00 15.13 1.00 15.32	B B	N C
ATOM	11632	CB	ASP	729	70. 146	40. 731	45.903	1.00 15.32	В	č
ATOM	11633	CG	ASP	729	70.034	42.019	46.696	1.00 18.11	В	C
ATOM	11634		ASP	729	69.663	43.055	46.104	1.00 20.57	В	0
ATOM ATOM	11635 11636	C C	ASP ASP	729 729	70. 317 72. 441	42.011 41.185	47. 907 45. 021	1.00 20.06 1.00 16.27	B B	0 C
ATOM	11637	ŏ	ASP	729	73. 253	40. 270	45.117	1.00 17.70	B	Ö
ATOM	11638	N	PHE	730	72. 772	42. 454	45. 211	1.00 16.74	В	N
ATOM ATOM	11639 11640	CA CB	PHE PHE	730 730	74. 136 75. 061	42. 824 42. 734	45.579 44.361	1.00 16.43 1.00 13.47	B B	C C
ATOM	11641	CG	PHE	730	74. 744	43. 728	43.304	1.00 13.41	В	. C
ATOM	11642	CD1	PHE	730	75. 282	45.006	43.355	1.00 12.64	В	C
ATOM	11643		PHE	730	73. 828	43. 423	42.303	1.00 12.46	В	C
ATOM ATOM	11644 11645		PHE PHE	730 730	74. 907 73. 446	45. 966 44. 377	42. 432 41. 376	1.00 11.61 1.00 9.11	B B	C
ATOM	11646		PHE	730	73. 986			1.00 10.39	В	č
ATOM	11647	C	PHE	730	74.112	44. 242	46.114	1.00 17.87	В	C
ATOM	11648	0	PHE	730	73. 094	44. 928	46.014 46.689	1.00 19.72	В	0
ATOM ATOM	11649 11650	N CA	GLN GLN	731 731	75. 230 75. 344	44. 673 46. 015	47. 246	1.00 18.41 1.00 17.25	B B	N C
ATOM	11651	CB	GLN	731	76.089	45. 961	48.569	1.00 18.02	B	č
ATOM	11652	CG	GLN	731	75. 547	44.948	49.536	1.00 25.59	В	C
ATOM ATOM	11653 11654	CD OF 1	GLN GLN	731 731	74. 087 73. 699	45. 183 46. 275	49. 854 50. 281	1.00 29.48 1.00 31.32	B B	C
ATOM	11655		GLN	731	73. 263	40. 273	49.647	1.00 31.32	В	O N
ATOM	11656	C	GLN	731	76. 124	46.889	46.272	1.00 16.69	B	Ç
MOTA	11657	0	GLN	731	77.060	46.417	45.623	1.00 13.71	В	0
ATOM ATOM	11658 11659	N CA	ALA ALA	732 732	75. 737 76. 425	48. 158 49. 084	46. 172 45. 284	1.00 15.59 1.00 15.79	B B	С
ATOM	11660	CB	ALA	732	75. 718	49. 147	43. 264	1.00 15.19	В	C
ATOM	11661	Č	ALA	732	76.540	50. 486	45.867	1.00 17.21	B	Ċ

					(Continued)
				FIG. 4 - 239	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11662 11663 11664 11665 11666 11667 11668 11669 11670 11671 11672 11673 11674	O ALA N MET CA MET CB MET CG MET SD MET CE MET C MET O MET N TRP CA TRP CB TRP	732 733 733 733 733 733 733 733 734 734 734	75. 769 50. 897 46. 734 1. 00 17. 93 B 77. 528 51. 220 45. 382 1. 00 17. 27 B 77. 737 52. 587 45. 812 1. 00 17. 39 B 78. 500 52. 628 47. 136 1. 00 18. 98 B 78. 775 54. 028 47. 661 1. 00 18. 20 B 77. 278 54. 979 47. 988 1. 00 21. 42 B 76. 781 54. 324 49. 578 1. 00 19. 12 B 78. 539 53. 268 44. 719 1. 00 17. 47 B 79. 604 52. 783 44. 318 1. 00 17. 30 B 78. 007 54. 378 44. 220 1. 00 16. 37 B 78. 673 55. 147 43. 175 1. 00 15. 48 B 77. 685 55. 428 42. 033 1. 00 14. 82 B 76. 691 56. 523 42. 353 1. 00 14. 06	O N C C C S C C O N C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11675 11676 11677 11678 11679 11680 11681 11682 11683 11684 11685	CD2 TRP CE2 TRP CE3 TRP CD1 TRP NE1 TRP CZ2 TRP CZ3 TRP CH2 TRP C TRP O TRP N TYR	734 734 734 734 734 734 734 734 734 735	75. 299 56. 363 42. 650 1. 00 12. 49 B 74. 785 57. 645 42. 939 1. 00 12. 15 B 74. 437 55. 259 42. 701 1. 00 12. 01 B 76. 953 57. 857 42. 468 1. 00 12. 61 B 75. 817 58. 535 42. 821 1. 00 13. 60 B 73. 449 57. 858 43. 276 1. 00 11. 75 B 73. 115 55. 466 43. 034 1. 00 13. 39 B 72. 629 56. 762 43. 319 1. 00 13. 13 B 79. 111 56. 457 43. 831 1. 00 13. 60 B 78. 491 56. 881 44. 788 1. 00 14. 71 B 80. 174 57. 090 43. 346 1. 00 13. 31 B	C C C C C C C C C C C C C C C C C C C
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11686 11687 11688 11689 11690 11691 11692 11693 11694 11695	CA TYR CB TYR CG TYR CD1 TYR CE1 TYR CD2 TYR CE2 TYR CZ TYR OH TYR C TYR	735 735	80. 598 58. 366 43. 926 1. 00 12. 17 B 81. 990 58. 260 44. 575 1. 00 10. 49 B 81. 964 57. 577 45. 920 1. 00 10. 18 B 81. 464 58. 232 47. 045 1. 00 11. 23 B 81. 321 57. 567 48. 272 1. 00 11. 72 B 82. 336 56. 241 46. 052 1. 00 11. 30 B 82. 198 55. 567 47. 270 1. 00 11. 75 B 81. 687 56. 235 48. 372 1. 00 12. 02 B 81. 511 55. 564 49. 563 1. 00 13. 79 B 80. 595 59. 430 42. 845 1. 00 14. 20 B	
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	11696 11697 11698 11699 11700 11701 11702 11703 11704	O TYR N THR CA THR CB THR OG1 THR CG2 THR C THR O THR N ASP	735 736 736 736 736 736 736 736 737	81. 391 59. 393 41. 910 1. 00 15. 56 B 79. 669 60. 372 42. 977 1. 00 15. 66 B 79. 517 61. 459 42. 026 1. 00 14. 01 B 78. 395 62. 401 42. 469 1. 00 13. 01 B 77. 163 61. 673 42. 534 1. 00 13. 00 B 78. 256 63. 571 41. 503 1. 00 11. 91 B 80. 789 62. 278 41. 882 1. 00 16. 80 B 81. 357 62. 730 42. 875 1. 00 19. 71 B 81. 230 62. 457 40. 640 1. 00 16. 82 B	O N C C O C C O N
ATOM ATOM ATOM ATOM ATOM ATOM	11705 11706 11707 11708 11709 11710	CA ASP CB ASP CG ASP OD1 ASP OD2 ASP C ASP	737 737 737 737 737 737	82. 407 63. 257 40. 322 1. 00 15. 22 B 82. 151 64. 728 40. 684 1. 00 15. 24 B 81. 101 65. 380 39. 785 1. 00 17. 61 B 80. 697 64. 753 38. 779 1. 00 16. 59 B 80. 680 66. 525 40. 078 1. 00 19. 23 B 83. 737 62. 811 40. 912 1. 00 15. 17 B	C C O O C

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ATOM 11755 O ALA 743 90.809 62.153 48.711 1.00 18.69 B O ATOM 11756 N SER 744 89.088 62.312 47.287 1.00 14.28 B N ATOM 11757 CA SER 744 88.681 63.556 47.908 1.00 14.62 B C											č
ATOM 11756 N SER 744 89.088 62.312 47.287 1.00 14.28 B N ATOM 11757 CA SER 744 88.681 63.556 47.908 1.00 14.62 B C											
ATOM 11757 CA SER 744 88.681 63.556 47.908 1.00 14.62 B C											
									1.00 14.62		С
	ATOM	11758	CB	SER	744	87. 369		47.321	1.00 16.50	В	С
ATOM 11759 OG SER 744 86.314 63.152 47.573 1.00 22.09 B 0	ATOM	11759	0G	SER	744	86. 314	63. 152	47. 573	1.00 22.09	В	0

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ATOM ATOM	11760 11761	C 0	SER SER	744 744	88. 515 88. 136		49. 390 49. 770	1.00 15.05 1.00 17.03	B B	C 0	
ATOM ATOM	11762 11763	N	SER	745	88. 822		50. 229	1.00 16.05	В	N	
ATOM	11764	CA CB	SER SER	745 745	88. 712 88. 811		51.666 52.361	1.00 15.38 1.00 15.23	B B	C	
ATOM	11765	0G	SER	745	88. 357	65.318	53.698	1.00 20.36	B	0	
ATOM	11766	C	SER	745	87. 427		52. 103	1.00 14.58	В	C	
ATOM ATOM	11767 11768	O N	SER THR	745 746	87. 467 86. 287		52. 773 51. 728	1.00 15.64 1.00 13.39	B B	O N	
ATOM	11769	CA	THR	746	85. 009		52. 121	1.00 13.33	В	C	
ATOM	11770	CB	THR	746	83. 836	64. 299	51.755	1.00 13.02	B	Č	
ATOM	11771		THR	746	83. 858	64. 579	50. 347	1.00 12.13	В	0	
ATOM ATOM	11772 11773	CG2	THR THR	746 746	83. 929		52. 547	1.00 6.36	В	C	
ATOM	11774	0	THR	746	84. 748 84. 382		51. 513 52. 215	1.00 13.71 1.00 13.77	B B	C 0	
ATOM	11775	N	ALA	747	84. 948		50. 211	1.00 15.70	В	N	
ATOM	11776	CA	ALA	747	84.698	60.575	49.556	1.00 17.75	. B	C	
ATOM ATOM	11777	CB	ALA	747	84.918		48. 047	1.00 18.85	В	C	
ATOM	11778 11779	C 0	ALA ALA	747 747	85. 579 85. 136		50. 133 · 50. 314	1.00 16.94 1.00 17.92	B B	C 0	
ATOM	11780	Ň	HIS	748	86. 828		50. 418	1.00 17.92	В	N	
ATOM	11781	CA	HIS	748	87.772	58. 873	50.987	1.00 15.53	B	Ċ	
ATOM	11782	CB	HIS	748	89.130		51. 194	1.00 14.50	В	C	
ATOM ATOM	11783 11784	CG CD2	HIS HIS	748 748	90. 106 90. 772		51.974 53.124	1.00 12.65	В	C	
ATOM	11785		HIS	748	90. 517		51.566	1.00 12.46 1.00 11.91	B B	C N	
ATOM	11786		HIS	748	91. 397		52.430	1.00 12.20	В	C	
ATOM	11787		HIS	748	91.569		53. 384	1.00 9.44	В	N	
ATOM ATOM	11788 11789	C 0	HIS HIS	748	87. 259		52. 316	1.00 15.00	В	C	•
ATOM	11790	N	GLN	748 749	87. 272 86. 808		52. 533 53. 200	1.00 14.52 1.00 14.63	B B	O N	•
ATOM	11791	CA	GLN	749	86. 283		54. 496	1.00 14.03	В	C	
ATOM	11792	CB	GLN	749	86.045	59.999	55.378	1.00 15.87	B	Č	
ATOM	11793	CC	GLN	749	87. 314	60. 722	55. 740	1.00 22.62	В	C .	
ATOM ATOM	11794 11795	CD OE1	GLN GLN	749 749	87. 056 86. 511	61.956 61.873	56. 564 57. 664	1.00 25.83 1.00 29.51	В	C	
ATOM	11796		GLN	749	87. 443	63. 116	56.039	1.00 25.51	B B	O N	
ATOM	11797	C	GLN	749	84. 984	57.999	54. 348	1.00 14.70	В	Ċ	
ATOM	11798	0	GLN	749	84. 749	57.015		1.00 14.10	В	0	
ATOM ATOM	11799 11800	N CA	HIS HIS	750 750	84. 147	58. 440	53. 415	1.00 13.44	В	N	
ATOM	11801		HIS	750 750	82. 865 82. 021	57. 808 58. 685	53. 174 52. 247	1.00 12.63 1.00 13.59	B B	.C	
ATOM	11802		HIS	750	80. 587	58. 272	52. 176	1.00 13.33	В	C	
ATOM	11803	CD2	HIS	750	79. 475	58.823	52. 713	1.00 13.33	B	Č	
ATOM	11804		HIS	750	80.175	57. 128	51.530	1.00 12.98	В	N	
ATOM .	11805 11806		HIS HIS	750 750	78. 869	56.992	51.673	1.00 14.44	В	C	
ATOM	11807	C	HIS	750 750	78. 419 82. 985	58.007 56.404	52.386 52.595	1.00 13.43 1.00 13.84	B B	N C	
ATOM	11808	Ŏ	HIS	750	82. 265	55. 499	53. 011	1.00 14.53	В	Ö	

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ATOM	11809	N	ILE	751	83. 885		51.638	1.00 13.03	В	N	
ATOM	11810	CA	ILE	751	84.013		51.077	1.00 12.47	В	C	
ATOM	11811	CB	ILE	751	84. 927		49. 814 50. 137	1.00 13.01	В	C	
ATOM ATOM	11812 11813	CG2	ILE ILE	751 751	86.326 84.999		49. 287	1.00 12.55 1.00 12.09	B B	C C C	
ATOM	11814	CD1	ILE	751	85. 677		47. 939	1.00 12.03	В	C	
ATOM	11815	CDI	ILE	751	84. 546		52.111	1.00 12.65	В	Č	
ATOM	11816	ŏ	ILE	751	84. 025		52. 241	1.00 12.49	B	ŏ.	
ATOM	11817	Ň	TYR	752	85.575		52.858	1.00 13.74	B	N	
ATOM	11818	CA	TYR	752	86.137	53.364	53.850	1.00 14.04	В	С	
ATOM	11819	CB	TYR	752	87.486		54.379	1.00 11.26	В	C	
ATOM	11820	CG	TYR	752	88.628		53.472	1.00 9.86	В	C	
ATOM	11821	CD1	TYR	752	89. 037		53.408	1.00 10.53	В	C	
ATOM	11822	CE1		752	90.015		52. 502	1.00 9.48	В	C	•
ATOM	11823		TYR	752	89. 235		52.608	1.00 9.66	В	C	
ATOM ATOM	11824 11825	CE2 CZ	TYR TYR	752 752	90. 219 90. 597		51.692 51.646	1.00 8.36 1.00 9.94	В	C	
ATOM	11826	OH	TYR	752	91.536		50. 739	1.00 9.94	B B	C 0	
ATOM	11827	C	TYR	752	85.170		54. 973	1.00 10.73	В	C	
ATOM	11828	ő	TYR	752	85. 176		55. 524	1.00 13.56	В	Ö	
ATOM	11829	Ň	THR	753	84. 323		55. 295	1.00 14.48	B	N	
ATOM	11830	CA	THR		83.316	53.864	56.330	1.00 14.27	B	Ċ	
ATOM	11831	CB	THR	753	82.582	55. 187	56.618	1.00 13.68	В	C	
ATOM	11832	0G1		753	83.519	56.136	57.130	1.00 17.48	В	0	
ATOM	11833	CG2		753	81.459	54. 987	57. 629	1.00 7.20	В	C	
ATOM	11834	C	THR	753	82. 301		55.815	1.00 16.15	В	C	
ATOM	11835	0	THR	753	81.958		56.508	1.00 18.93	В	0	
ATOM	11836	N	HIS	754	81.830		54. 589	1.00 15.38	В	N	
ATOM	11837	CA	HIS	754	80.840		53.999	1.00 16.06	В	C	
ATOM ATOM	11838 11839	CB CG	HIS HIS	754 754	80. 424 79. 109		52.620 52.162	1.00 15.26	В	C	
ATOM	11840		HIS	754 754	78. 779		51.095	1.00 16.39 1.00 15.75	B B	C C	
ATOM	11841		HIS	754	77. 936		52.850	1.00 13.13	В	N	
ATOM	11842		HIS	754	76.940			1.00 15.86	В	Č	
ATOM	11843		HIS	754	77. 425		51.161	1.00 17.13	В	Ň	
ATOM	11844	C	HIS	754	81.349		53.886	1.00 16.28	B	Ċ	
ATOM	11845	0	HIS	754	80.639		54.238	1.00 17.31	В	Ō	
ATOM	11846	N	MET	755	82.571	50.564	53.383	1.00 15.98	В	N	
ATOM	11847	CA	MET	7 55	83. 158		53. 250	1.00 16.05	В	C	
ATOM	11848	CB	MET	755	84. 532		52. 573	1.00 15.41	В	C	
ATOM	11849	- CG	MET	755	84. 491	49.542	51.081	1.00 17.11	В	C	
ATOM	11850	SD	MET	755 755	86.112		50. 322	1.00 18.41	В	S	
ATOM	11851	CE	MET	755 755	86.882		50.742	1.00 20.74	В	. C	
ATOM ATOM	11852 11853	C 0	MET MET	755 755	83. 309 83. 080		54. 623 54. 783	1.00 15.38 1.00 13.30	B B	C	
ATOM	11854	N	SER	756	83. 701	49.371	54. 783 55. 614	1.00 13.30	В	O N	
ATOM	11855	CA	SER	756	83. 854		56.946	1.00 18.52	В	C	
ATOM	11856	CB	SER	756	84. 413		57. 878	1.00 18.88	В	Č	
ATOM	11857	0G	SER	756	85. 723		57. 477	1.00 18.74	B	ŏ	

		(Continued)
	FIG. 4-243	
ATOM 11859 O SER 7 ATOM 11860 N HIS 7 ATOM 11861 CA HIS 7 ATOM 11862 CB HIS 7 ATOM 11863 CG HIS 7 ATOM 11864 CD2 HIS 7 ATOM 11865 ND1 HIS 7 ATOM 11866 CE1 HIS 7 ATOM 11866 CE1 HIS 7 ATOM 11867 NE2 HIS 7 ATOM 11868 C HIS 7 ATOM 11869 O HIS 7 ATOM 11870 N PHE 7 ATOM 11871 CA PHE 7 ATOM 11872 CB PHE 7 ATOM 11873 CG PHE 7 ATOM 11874 CD1 PHE 7 ATOM 11875 CD2 PHE 7 ATOM 11876 CE1 PHE 7 ATOM 11876 CE1 PHE 7 ATOM 11877 CE2 PHE 7 ATOM 11878 CZ PHE 7 ATOM 11878 CZ PHE 7 ATOM 11879 C PHE 7 ATOM 11880 O PHE 7 ATOM 11881 N ILE 7 ATOM 11882 CA ILE 7 ATOM 11884 CG2 ILE 7 ATOM 11885 CG1 ILE 7 ATOM 11886 CD1 ILE 7 ATOM 11887 C ILE 7 ATOM 11887 C ILE 7 ATOM 11888 O ILE 7 ATOM 11889 N IVS	58 80. 941 44. 120 52. 254 1. 00 6 58 78. 334 43. 716 51. 409 1. 00 9 58 80. 638 43. 045 51. 417 1. 00 6 58 79. 340 42. 836 50. 991 1. 00 2 58 80. 697 44. 674 55. 560 1. 00 20 58 80. 110 43. 631 55. 851 1. 00 21 59 82. 014 44. 811 55. 654 1. 00 23 59 82. 858 43. 722 56. 117 1. 00 25 59 84. 364 44. 129 56. 069 1. 00 25 59 84. 994 44. 041 57. 437 1. 00 26 59 84. 706 43. 263 53. 704 1. 00 26 59 84. 706 43. 263 53. 704 1. 00 26 59 82. 441 43. 318 57. 529 1. 00 25 59 82. 420 42. 136 57. 866 1. 00 26	1. 14 B C 1. 94 B O 1. 68 B N 1. 20 B C 1. 13 B C 1. 10 B C 1. 62 B N 1. 60 B C 1. 81 B N 1. 94 B C 1. 58 B O 1. 70 B C 1. 77 B C 1. 77 B C 1. 77 B C 1. 77 B C 1. 78
ATOM 11890 CA LYS 76 ATOM 11891 CB LYS 76 ATOM 11892 CG LYS 76	60 81.671 44.012 59.713 1.00 26. 60 81.444 45.300 60.487 1.00 26. 60 82.178 45.298 61.792 1.00 29.	B C 43 B C .00 B C
ATOM 11894 CE LYS 76 ATOM 11895 NZ LYS 76 ATOM 11896 C LYS 76	60 83.666 45.271 61.537 1.00 28. 60 84.139 46.665 61.250 1.00 30. 60 83.776 47.523 62.420 1.00 31. 60 80.406 43.179 59.740 1.00 27. 60 80.312 42.200 60.473 1.00 28.	B C 29 B N 08 B C
ATOM 11898 N GLN 76 ATOM 11899 CA GLN 76 ATOM 11900 CB GLN 76 ATOM 11901 CG GLN 76	61 79. 431 43. 581 58. 940 1. 00 28. 61 78. 170 42. 866 58. 844 1. 00 29. 61 77. 213 43. 652 57. 942 1. 00 31. 61 76. 072 42. 855 57. 347 1. 00 34.	08 B N 69 B C 26 B C 99 B C
ATOM 11902 CD GLN 76 ATOM 11903 OE1 GLN 76 ATOM 11904 NE2 GLN 76 ATOM 11905 C GLN 76 ATOM 11906 O GLN 76	61 76.800 42.775 55.062 1.00 37. 61 76.464 40.808 56.112 1.00 39. 61 78.401 41.456 58.295 1.00 30.	29 B O 80 B N 00 B C

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(Continued) FIG. 4-244 57.320 1.00 29.71 N 41.333 79.291 11907 N CYS 762 ATOM C 1.00 30.30 79.588 40.035 56.731 CYS 762 11908 CA ATOM C 57.712 1.00 30.21 В 80.275 39,077 CYS 762 ATOM 11909 C 0 37.860 57.578 1.00 29.67 В 80.153 11910 CYS 762 **ATOM** 0 1.00 30.01 В C 40.212 55.474 **CYS** 762 80.458 11911 CB **ATOM** В S 38.665 54.849 1.00 33.72 **CYS** 81.198 SG 762 **ATOM** 11912 1.00 30.53 В N 58.698 80.986 39.618 N PHE 763 11913 **ATOM** C B 59.664 1.00 31.28 81.694 38.783 PHE 763 ATOM 11914 CA C В 39.310 59.885 1.00 29.29 763 83.112 CB PHE ATOM 11915 Ċ 58.736 1.00 27.21 В 763 84.052 39.057 CG PHE ATOM 11916 38.280 57.650 1.00 26.19 C В 763 83.663 CD1 PHE ATOM 11917 Ċ 85.348 39.572 58.762 1.00 26.38 B ATOM 11918 CD2 PHE 763 C В 56.605 1.00 27.91 11919 CE1 PHE 763 84.552 38.015 ATOM C 57.727 В 86.249 39.316 1.00 27.36 CE2 PHE 763 11920 ATOM C 38.533 56.643 1.00 27.55 В 85.851 CZPHE 763 ATOM 11921 C В 61.011 1.00 34.52 80.994 38.666 ATOM 11922 C PHE 763 В 0 763 81.473 37.970 61.908 1.00 32.78 11923 0 PHE ATOM 79.862 39.346 61.151 1.00 39.49 В N 764 ATOM 11924 N SER 62.393 39.319 1.00 43.60 C 79.099 В 11925 CA SER 764 ATOM 62.273 В C 77.860 40.199 1.00 44.56 764 ATOM 11926 CB SER 61.948 В 0 78.218 41.528 1.00 50.05 **ATOM** 11927 SER 764 0G 62.746 1.00 45.96 78.668 37.909 В C SER **ATOM** 11928 C 764 77.885 62.028 1.00 45.86 В 0 11929 0 SER 764 37.289 **ATOM** 79.189 37.404 63.856 1.00 49.22 В N **ATOM** 11930 N LEU 765 C 36.070 64.317 1.00 52.03 В 11931 CA LEU 765 78.845 **ATOM** 79.754 765 35.678 65.481 1.00 52.53 В C 11932 CB LEU ATOM 81.234 35.558 65.115 1.00 52.85 В C 11933 CG LEU 765 ATOM 35.452 66.376 1.00 53.55 В C 82.074 **ATOM** 11934 CD1 LEU 765 В C 11935 CD2 LEU 765 81.435 34.344 64. 214 1.00 52.54 ATOM 77.383 36.069 64.761 1.00 54.34 В C **ATOM** 11936 C LEU 765 65.743 1.00 53.63 0 0 LEU 765 77.019 36.721 В 11937 ATOM 35.340 64.031 1.00 56.38 В N 76.523 ATOM 11938 N PR₀ 766 76.833 34.541 62.831 1.00 56.67 В C 11939 CD PR₀ 766 ATOM 75.095 35.263 64.356 1.00 57.95 C В 766 ATOM 11940 CA PR0 PR₀ 766 74.509 34.544 63.141 1.00 58.24 В 11941 CB ATOM 62.728 1.00 57.40 В 11942 CG PR₀ 766 75.626 33.633 ATOM 11943 74.805 34.523 65.664 1.00 59.30 В C C **PRO** 766 ATOM 65.711 1.00 60.29 73.791 33.789 В 0 11944 PR₀ 766 ATOM 0 11945 75.584 34.704 66.627 1.00 59.84 В OXT PRO ATOM 766 TER 11946 PR₀ 766 11947 C1 NAG 901 25.105 38.477 14.927 1.00 45.03 E C ATOM 26.266 38.501 13.922 1.00 45.16 E 11948 C2NAG 901 ATOM 27.447 14.595 1.00 44.20 E 39.002 N NAG 901 ATOM 11949 N2 28.662 38.702 14.153 1.00 43.63 E C 11950 **C7** NAG 901 ATOM 29.050 13.997 1.00 44.60 37.546 E 0 **ATOM** 11951 07 NAG 901 **C8** NAG 901 29.588 39.864 13.838 1.00 43.83 E C ATOM 11952 25.942 12.713 1.00 46.38 C C3NAG 901 39.385 11953 ATOM 1.00 49.49 901 26.953 39.235 11.728 E NAG 0 03 ATOM 11954

38.987 SUBSTITUTE SHEET (RULE 26)

12.124

1.00 47.76

E

24. 591

901

ATOM

11955

C4

NAG

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					EIC 4-245	(Continued)
					FIG. 4 - 245	
ATOM ATOM	11956 11957	04 C5	NAG NAG	901 901	24. 256 39. 836 11. 036 1. 00 49. 01 E 23. 545 39. 104 13. 219 1. 00 49. 11 E	0 C
ATOM	11958	05	NAG	901	23. 858 38. 173 14. 276 1. 00 47. 99 E	0
ATOM	11959	C6	NAG	901	22. 143 38. 804 12. 731 1. 00 50. 99 E	C
ATOM	11960	06	NAG	901	21. 706 39. 781 11. 793 1. 00 53. 28 E 34. 526 67. 450 4. 248 1. 00 29. 71 E	0
ATOM ATOM	11961 11962	C1 C2	NAG NAG	902 902	34.526 67.450 4.248 1.00 29.71 E 33.682 66.990 3.051 1.00 31.02 E	C C
ATOM	11963	N2	NAG	902	34. 077 65. 638 2. 692 1. 00 35. 02 E	Ň
ATOM	11964	C7	NAG	902	33.181 64.660 2.610 1.00 35.78 E	С
ATOM	11965	07	NAG	902	32. 213 64. 701 1. 852 1. 00 37. 59 E	0
ATOM	11966	C8	NAG	902	33. 392 63. 449 3. 503 1. 00 37. 18 E 33. 927 67. 915 1. 848 1. 00 31. 67 E	C C
ATOM ATOM	11967 11968	C3 03	NAG NAG	902 902	33. 927 67. 915 1. 848 1. 00 31. 67 E 33. 032 67. 583 0. 794 1. 00 34. 76 E	0
ATOM	11969	C4	NAG	902	33. 753 69. 386 2. 248 1. 00 31. 76 E	č
ATOM	11970	04	NAG	902	34. 037 70. 238 1. 144 1. 00 30. 03 E	0
ATOM	11971	C5	NAG	902	34. 701 69. 674 3. 412 1. 00 30. 64 E	C .
ATOM ATOM	11972 11973	05 C6	NAG	902 902	34. 332 68. 844 4. 526 1. 00 30. 02 E 34. 720 71. 114 3. 892 1. 00 30. 81 E	0
ATOM	11973	06	NAG NAG	902	34.720 71.114 3.892 1.00 30.81 E 33.457 71.512 4.409 1.00 34.26 E	C 0
ATOM	11975	CI	NAG	903	64. 239 77. 734 14. 341 1. 00 27. 20 E	Č
ATOM	11976	C2	NAG	903	63. 984 78. 203 12. 917 1. 00 26. 96 E	C
ATOM	11977	N2	NAG	903	63. 551 77. 080 12. 116 1. 00 25. 19 E	N
ATOM ATOM	11978 11979	C7 07	NAG NAG	903 903	62.349 77.076 11.551 1.00 24.99 E 62.121 76.492 10.490 1.00 25.88 E	C
ATOM	11980	C8	NAG	903	62.121 76.492 10.490 1.00 25.88 E 61.222 77.800 12.272 1.00 23.55 E	0 C
ATOM	11981	C3	NAG	903	65. 253 78. 817 12. 325 1. 00 29. 00 E	č
ATOM	11982	03	NAG	903	64. 947 79. 400 11. 066 1. 00 29. 62 E	0
ATOM	11983	C4	NAG	903	65. 814 79. 900 13. 248 1. 00 30. 83 E	C
ATOM ATOM	11984 11985	04 C5	NAG NAG	903 903	67. 092 80. 316 12. 778 1. 00 31. 15 E 65. 929 79. 389 14. 690 1. 00 30. 71 E	0
ATOM	11986	05	NAG	903	65. 929 79. 389 14. 690 1. 00 30. 71 E 64. 669 78. 842 15. 133 1. 00 30. 11 E	C 0
ATOM	11987	C6	NAG	903	66. 276 80. 502 15. 659 1. 00 32. 26 E	č
ATOM	11988	06	NAG	903	65. 937 80. 144 16. 993 1. 00 35. 52 E	0 .
ATOM	11989	C1	NAG	904	56. 857 73. 229 -0. 933 1. 00 21. 65 E	C
ATOM ATOM	11990 11991	C2 N2	NAG NAG	904 904	58. 289 73. 099 -1. 475 1. 00 21. 59 E 58. 532 71. 758 -1. 961 1. 00 21. 40 E	C
ATOM	11991	C7	NAG	904	58.532 71.758 -1.961 1.00 21.40 E 58.567 71.523 -3.267 1.00 20.76 E	N C
ATOM	11993	07	NAG	904	58. 745 72. 412 -4. 104 1. 00 18. 55 E	ŏ
ATOM	11994	C8	NAG	904	58.371 70.080 -3.709 1.00 20.74 E	C
ATOM	11995	C3	NAG	904		C
ATOM	11996	03	NAG	904	60.611 73.413 -1.009 1.00 22.81 E	0
ATOM ATOM	11997 11998	C4 04	NAG NAG	904 904	59. 022 74. 832 0. 129 1. 00 22. 85 E 59. 986 75. 217 1. 101 1. 00 24. 62 E	C 0
ATOM	11999	C5	NAG	904	57.634 74.781 0.737 1.00 22.86 E	C
ATOM	12000	05	NAG	904	56.672 74.506 -0.297 1.00 21.95 E	ŏ
ATOM	12001	C6	NAG	904	57. 232 76. 083 1. 385 1. 00 24. 39 E	C
ATOM	12002	06	NAG	904	57.196 77.133 0.430 1.00 31.81 E	0
ATOM ATOM	12003 12004	C1 C2	NAG NAG	905 905	49.743 85.075 37.084 1.00 31.93 E	C
VION	12004	U4	וותט	900	49.010 86.230 37.756 1.00 33.35 E	C

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					Te T	C 1	216			(Continued)
					FI	G. 4 -	240			
ATOM	12005	N2	NAG	905	47.823	86. 586	37.012	1.00 34.30	Е	N
ATOM	12006	C7	NAG	905	46.648	86.099	37. 395	1.00 35.18	E	C
ATOM	12007	07	NAG	905	46. 362		38. 578	1.00 36.47	Е	0
ATOM	12008	C8	NAG	905	45.640		36. 303	1.00 37.15	Е	C
ATOM	12009	C3	NAG	905	49.951		37.924	1.00 33.45	E	C
ATOM	12010	03	NAG	905	49. 256	88.512	38.495	1.00 33.93	Е	0
ATOM	12011	C4	NAG	905	51.043		38.863	1.00 35.37	E	C
ATOM	12012	04	NAG	905	51.934		39.193	1.00 35.45	E	0
ATOM	12013	C5	NAG	905	51.794	85.773	38. 215	1.00 34.39	Е	C
ATOM	12014	05	NAG	905	50.878	84.684	37.887	1.00 32.56	E	0
ATOM	12015	C6	NAG	905	52. 787	85. 212	39. 214	1.00 36.29	E	C
ATOM	12016	06	NAG	905	52.150	84. 936	40.459	1.00 35.52	E	0
ATOM	12017	C1	NAG	906	128.439	74.792	56.371	1.00 36.45	E	C
ATOM	12018	C2	NAG	906	127. 977	75.856	55.375	1.00 37.00	E	C
ATOM	12019	N2	NAG	906	126.880	75.335	54.586	1.00 37.17	E	N
ATOM	12020	C7	NAG	906	125.666	75.871	54.690	1.00 38.41	Е	C
ATOM	12021	07	NAG	906	125. 264		55.714	1.00 38.52	Е	0
ATOM	12022	C8	NAG	906	124.760		53. 471	1.00 36.25	Е	C
ATOM	12023	C3	NAG	906	129. 133		54.465	1.00 38.66	Е	C
ATOM	12024	03	NAG	906	128. 723		53.625	1.00 39.59	Е	. 0
ATOM	12025	C4	NAG	906	130. 331		55.308	1.00 39.58	E	C
ATOM	12026	04	NAG	906	131.439		54.460	1.00 41.48	E	0
ATOM	12027	C5	NAG	906	130. 699		56.312	1.00 40.24	E	C
ATOM	12028	05	NAG	906	129. 556		57. 133	1.00 38.27	E	0
ATOM	12029	C6	NAG	906	131.811	76.032	57. 255	1.00 41.89	E	C
ATOM	12030	06	NAG	906	131.906		58.378	1.00 46.70	E	0
ATOM	12031	C1	NAG	907	126. 770		25.405	1.00 33.54	E	C
ATOM	12032	C2	NAG	907	127. 763		25.478	1.00 35.73	Е	C
ATOM	12033	N2	NAG	907	127. 401	74. 367	26.540	1.00 37.97	E	N
ATOM	12034	C7	NAG	907	128. 139		27.644	1.00 41.34	E	C
ATOM	12035	07	NAG	907	128. 715	73. 403	28.094	1.00 42.96	E	0
ATOM	12036	C8	NAG	907	128. 278	75. 739	28. 352	1.00 42.60	E	C
ATOM	12037	C3	NAG	907	127. 776	74. 167	24. 126	1.00 36.63	E	C
ATOM	12038	03	NAG	907	128. 692	75. 253	24. 154	1.00 38.28	E	0
ATOM	12039	C4	NAG	907	128. 171	73. 148	23.047	1.00 35.89	E	C
ATOM	12040	04	NAG	907	128. 191	73. 758	21.763	1.00 35.82	E	0
ATOM	12041	C5	NAG	907	127. 161	71.995	23.075	1.00 35.12	E	C
ATOM	12042	05	NAG	907	127. 166	71.377	24.380	1.00 32.61	E	0
ATOM	12043	C6	NAG	907	127. 444	70.913	22.057	1.00 36.17	E	С
ATOM	12044	06	NAG	907	128. 515	70.083	22.478	1.00 38.44	E	0
ATOM	12045	C1	NAG	908	97.567	64. 129	12.586	1.00 33.83	E	C
ATOM	12046	C2	NAG	908	98. 226	65. 101	11.602	1.00 36.51	E	С
ATOM	12047	N2	NAG	908	98. 466	66.365	12. 269	1.00 40.33	E	Ŋ
ATOM	12048	C7	NAG	908	99.645	66. 962	12.148	1.00 43.03	Ē	C
ATOM	12049	07	NAG	908	100. 703	66. 434	12.500	1.00 45.77	E	0
ATOM	12050	C8	NAG	908	99.655	68. 349	11.529	1.00 43.86	E	C
ATOM	12051	C3	NAG	908	97. 328	65. 325	10.380	1.00 37.11	E	C
ATOM	12052	03	NAG	908	98. 013	66. 122	9. 426	1.00 37.35	E	0
ATOM	12053	C4	NAG	908	96.945	63. 975	9.760	1.00 36.97	Е	C

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										(Conti	nued)
					FIG	}. 4 -	2 4 7			, • •	
ATOM	12054	04	NAG	908	96.049	64. 165	8. 668	1.00 36.08	Е	0	
ATOM	12055	C5	NAG	908	96. 291	63.106	10.841	1.00 35.43	Ē	Č	
ATOM	12056	05	NAG	908	97.215	62.906	11.930	1.00 33.34	Ē	Ŏ	
ATOM	1.2057	C6	NAG	908	95.890	61.735	10.341	1.00 36.72	Ē	Č	
ATOM	12058	06	NAG	908	95.085	61.057	11.296	1.00 38.75	Ē	0	•
ATOM	12059	C1	NAG	909	106.501	80.407	11.987	1.00 55.21	Ē	Č	
ATOM	12060	C2	NAG	909	105.627	81.255	11.048	1.00 55.75	Ē	Č	
ATOM	12061	N2	NAG	. 909	105.631	82.658	11.427	1.00 55.80	Ē	N	
ATOM	12062	C7	NAG	909	106.748	83.259	11.828	1.00 56.83	E	C	
ATOM	12063	07	NAG	909	107.685	83.526	11.066	1.00 55.16	E	0	
ATOM	12064	C8	NAG	909	106.838	83.620	13.305	1.00 56.25	E	C	
ATOM	12065	C3	NAG	909	104.195	80.724	11.087	1.00 56.36	E	С	
ATOM	12066	03	NAG	909	103.396	81.452	10.166	1.00 58.58	E	0	
ATOM	12067	C4	NAG	909	104.176	79. 229	10.744	1.00 56.19	Е	C	
ATOM	12068	04	NAG	909	102.855	78.716	10.862	1.00 55.29	Е	0	
ATOM	12069	C5	NAG	909	105.117	78.478	11.692	1.00 56.24	E	C	
ATOM	12070	05	NAG	909	106.446	79.028	11.600	1.00 56.65	Е	0	
ATOM	12071	C6	NAG	909	105.230	76.996	11.381	1.00 57.38	E	C	
ATOM	12072	06	NAG	909	106.370	76.423	12.010	1.00 55.01	Е	0	
ATOM	12073	C1	NAG	910	105. 213	38. 428	20.006	1.00 34.33	Е	C	
ATOM	12074	C2	NAG	910	106.113	37. 293	19.498	1.00 37.27	E	C	
ATOM	12075	N2	NAG	910	107.447	37. 789	19. 211	1.00 40.05	E	N	
ATOM	12076	C7	NAG	910	108. 495	36. 984	19.368	1.00 42.24	E	C	
ATOM	12077	07	NAG	910	109.013	36. 771	20. 465	1.00 42.65	E	0	
ATOM	12078	C8	NAG	910	109.047	36. 295	18. 126	1.00 42.65	E	C	
ATOM	12079	C3	NAG	910	105.504	36.650	18. 245	1.00 37.60	E	C	
ATOM	12080	03	NAG	910	106. 296	35. 547	17. 831	1.00 38.44	E	0	
ATOM	12081	C4	NAG	910	104.084	36. 182	18. 551	1.00 36.63	E	C	
ATOM	12082	04	NAG	910	103. 489	35.616	17. 388	1.00 37.52	E	0	
ATOM	12083	C5	NAG	910	103. 274	37. 387	19.037	1.00 35.81	E	C	
ATOM	12084	05	NAG	910	103.883	37. 930	20. 229	1.00 34.96	E	0	
ATOM	12085	C6	NAG	910	101.838	37. 042	19. 385	1.00 34.79	Ē	C	
ATOM TER	12086	06	NAG	910	101.781	36. 089	20. 437	1.00 34.77	E	0	
ATOM	12087 12088	٥	NAG	910	E9 49E	00 704	10 170	1 00 10 00	E		
ATOM	12089	0	HOH	1	53. 435	80. 704	18. 172	1.00 10.60	W	0	
ATOM	12009	0	HOH HOH	2	57.473	78. 703	26. 320	1.00 21.03	W	0	
ATOM	12090	0	НОН	3	65.386	56.077	37.040	1.00 7.09	W	0	
ATOM	12092	0	НОН	4 5	56.235 58.127	76. 520	22. 816 28. 066	1.00 14.76	W	0	
ATOM	12093	0	НОН	6	40.099	60.758		1.00 4.57	W	0	
ATOM	12094	0	НОН	7	29. 796	59. 877 47. 323	48. 410	1.00 16.00	W	0	
ATOM	12095	0	HOH	8			37. 410	1.00 24.76	W	0	
ATOM	12096	0	нон НОН	9	38.634 41.732	67. 195 52. 103	51. 371 37. 673	1.00 22.65	W	0	
ATOM	12097	0	НОН	10	79. 275	54. 159	21. 409	1.00 13.34	W	0	
ATOM	12098	0	HOH	11	65. 287	66. 160	35. 128	1.00 15.53 1.00 7.29	W	0	
ATOM	12099	0	HOH	12	79. 267	49. 364	26. 780	1.00 7.29 1.00 14.00	W	0	
ATOM	12100	Ö	HOH	13	67. 989	56. 792	26. 833	1.00 14.00	n Y	0 0	
ATOM	12101	Ö	НОН	14.	68.995	70. 138	19.815	1.00 20.21	W	0	
ATOM	12102	ŏ	НОН	15	59. 193	63. 441	21. 787	1.00 12.38	"	0	
		•			100	x 1 1	21. (0)	** 00 0. 00	11	v	

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					FIG. 4-248	(Continued)
ATOM	12103	0	НОН	16	49.896 66.700 47.886 1.00 13.21 W	0
ATOM	12104	0	НОН	17	48. 544 53. 043 50. 567 1. 00 20. 65 W	0
ATOM	12105	0	HOH	18	73. 938 69. 817 · 52. 424 1. 00 34. 74 W	0
ATOM	12106	0	HOH	19	36. 883 69. 650 29. 378 1. 00 25. 18 W	0
ATOM	12107	0	HOH	20	50. 912 61. 115 48. 431 1. 00 18. 77 W	0
ATOM ATOM	12108 12109	0	НОН НОН	21 22	58. 369 85. 282 28. 107 1. 00 27. 06 W 62. 886 63. 930 21. 686 1. 00 29. 16 W	0
ATOM	12110	0	ноп НОН	23		0
ATOM	12111	0	НОН	23 24	43.777 87.394 23.730 1.00 9.96 W 48.078 67.109 30.405 1.00 21.66 W	0
ATOM	12112	ŏ	НОН	25	36. 753 80. 303 31. 025 1. 00 34. 33 W	0
ATOM	12113	ŏ	НОН	26	63. 225 66. 634 22. 568 1. 00 10. 18 W	0
ATOM	12114	Õ	НОН	$\frac{27}{27}$	35. 078 54. 838 52. 427 1. 00 29. 90 W	ŏ
ATOM	12115	Ŏ	НОН	28	57. 184 80. 961 23. 145 1. 00 17. 51 W	ŏ
ATOM	12116	0	HOH	29	73.677 71.484 27.824 1.00 34.92 W	Ŏ
ATOM	12117	0	HOH	30	76. 251 57. 060 34. 794 1. 00 28. 05 W	0
ATOM	12118	0	HOH	31	72. 985 72. 092 24. 987 1. 00 14. 46 W	0
ATOM	12119	0	HOH	32	61. 839 84. 543 25. 502 1. 00 22. 75 W	0
ATOM	12120	0	HOH	33	33. 787 63. 840 46. 551 1. 00 12. 55 W	0
ATOM	12121	0	HOH	34	47. 827 47. 441 47. 587 1. 00 25. 33 W	0
ATOM	12122	0	НОН	35	55. 562 56. 510 44. 904 1. 00 30. 51 W	0
ATOM	12123	0	НОН	36	31. 114 59. 222 42. 224 1. 00 13. 22 W	0
ATOM	12124	0	HOH	37	82. 143 64. 199 47. 510 1. 00 21. 69 W	0
ATOM	12125	0	НОН	38	41. 587 70. 385 33. 904 1. 00 24. 19 W	0
ATOM ATOM	12126 12127	0	НОН НОН	39 40	70. 447 47. 056 34. 998 1. 00 24. 19	0
ATOM	12128	0	НОН	40 41	23. 146 49. 571 32. 910 1. 00 22. 85 W 23. 427 53. 516 39. 573 1. 00 12. 47 W	0
ATOM	12129	ő	HOH	42		0
ATOM	12130	ŏ	НОН	43	74.977 48.248 21.021 1.00 24.35 W 81.171 53.457 19.457 1.00 32.23 W	0 0
ATOM	12131	ŏ	НОН	44	70. 982 61. 003 21. 232 1. 00 19. 07	0
ATOM	12132	Õ.	НОН	45	51.713 50.325 19.619 1.00 36.05 W	0
ATOM	12133	0	НОН	46	75. 424 58. 001 59. 062 1. 00 20. 53	Ŏ
ATOM	12134	0	HOH	47	52. 251 54. 978 15. 598 1. 00 20. 74 W	Ŏ
ATOM	12135	0	HOH	48	37. 551 51. 103 23. 882 1. 00 16. 65 W	Ö
ATOM	12136	0 -	HOH	49	31.428 66.281 21.097 1.00 18.82 W	0
	12137			50 -	45.546 72.589 -9.525 1.00 19.51 W	0
ATOM	12138	0	HOH	51	71.765 47.337 39.374 1.00 16.49 W	0
ATOM	12139	0	НОН	52	57.328 68.673 61.331 1.00 26.41 W	0
ATOM	12140	0	НОН	53	72. 778 48. 947 47. 621 1. 00 17. 49 W	0
ATOM	12141	0	HOH	54	30. 292 82. 021 10. 956 1. 00 24. 56 W	0
ATOM	12142	0	HOH	55 56	47. 165 45. 427 40. 043 1. 00 35. 52 W	0
ATOM	12143	0	HOH	56	25. 673 60. 491 43. 209 1. 00 10. 79 W	0
ATOM ATOM	12144 12145	0	НОН НОН	57 58	71. 617 62. 843 34. 752 1. 00 17. 19 W	0
ATOM	12145	0	HOH	58 59	46.059 55.643 2.123 1.00 19.51 W 68.766 45.985 50.017 1.00 22.18 W	0
ATOM	12140	0	НОН	60		0 ·
ATOM	12148	Ŏ	НОН	61	52.732 70.566 0.317 1.00 32.17 W 61.782 69.597 25.094 1.00 13.27 W	0 0
ATOM	12149	Ö	НОН	62	51. 352 79. 521 14. 538 1. 00 17. 25	0
ATOM	12150	ŏ	НОН	63	48. 267 86. 907 16. 122 1. 00 21. 54	0
ATOM	12151	Ŏ	НОН	64	49. 536 54. 337 14. 938 1. 00 22. 27 W	0
					11,000 81,00 88181 11	•

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					FIG. 4-249	(Continued)
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	12152 12153 12154 12155 12156 12157 12158 12159 12160	0 0 0 0 0 0 0 0	HOH HOH HOH HOH HOH HOH HOH	65 66 67 68 69 70 71 72 73	37. 711 84. 458 31. 782 1. 00 38. 65 W 41. 832 62. 441 48. 190 1. 00 23. 50 W 56. 514 63. 214 39. 402 1. 00 20. 39 W 48. 166 60. 456 42. 122 1. 00 37. 55 W 52. 076 51. 584 45. 757 1. 00 22. 02 W 47. 607 61. 634 15. 612 1. 00 34. 50 W 39. 108 76. 636 34. 882 1. 00 24. 21 W 62. 894 85. 163 44. 724 1. 00 38. 05 W 49. 937 51. 963 48. 658 1. 00 25. 50 W	0 0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	12161 12162 12163 12164 12165 12166 12167 12168	0 0 0 0 0 0 0 0	HOH HOH HOH HOH HOH HOH HOH	74 75 76 77 78 79 80 81	32. 972 63. 405 9. 645 1. 00 31. 16 W 76. 481 50. 940 55. 523 1. 00 8. 02 W 54. 751 68. 666 -3. 038 1. 00 19. 33 W 69. 797 76. 851 37. 550 1. 00 38. 44 W 60. 195 69. 793 56. 043 1. 00 27. 75 W 68. 721 77. 775 28. 423 1. 00 14. 61 W 76. 538 41. 044 29. 727 1. 00 24. 17 W	0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM	12169 12170 12171 12172 12173 12174 12175	0 0 0 0 0 0	HOH HOH HOH HOH HOH HOH	82 83 84 85 86 87 88	42. 573 57. 621 42. 066 1. 00 19. 56 W 51. 219 56. 139 24. 829 1. 00 41. 31 W 64. 281 54. 295 25. 797 1. 00 15. 83 W 48. 093 54. 052 46. 307 1. 00 38. 41 W 37. 006 52. 225 21. 202 1. 00 23. 83 W 44. 149 74. 948 5. 314 1. 00 17. 55 W 72. 912 75. 091 28. 633 1. 00 25. 98 W	0 0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	12176 12177 12178 12179 12180 12181 12182 12183	0 0 0 0 0 0	HOH HOH HOH HOH HOH HOH HOH	89 90 91 92 93 94 95	52. 329 67. 860 33. 481 1.00 8.31 W 66. 266 74. 773 42. 238 1.00 16.00 W 59. 283 77. 076 9. 072 1.00 41. 29 W 77. 526 46. 454 20. 254 1.00 34. 51 W 59. 751 56. 673 29. 191 1.00 24. 40 W 43. 531 63. 248 14. 122 1.00 22. 64 W 56. 677 73. 257 -8. 550 1.00 18. 65 W 64. 366 82. 016 33. 202 1.00 24. 81 W	0 0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	12184 12185 12186 12187 12188 12189 12190 12191	0 0 0 0 0 0	HOH HOH HOH HOH HOH HOH HOH	97 98 99 100 101 102 103 104	58. 839 62. 776 26. 537 1. 00 11. 00 W 52. 478 72. 152 3. 092 1. 00 13. 58 W 59. 860 59. 389 29. 429 1. 00 20. 06 W 64. 047 73. 184 44. 557 1. 00 15. 66 W 44. 369 74. 978 38. 087 1. 00 11. 11 W 61. 861 50. 833 14. 510 1. 00 31. 09 W 40. 708 73. 940 22. 137 1. 00 13. 81 W	0 0 0 0 0 0
ATOM ATOM ATOM ATOM ATOM ATOM ATOM ATOM	12192 12193 12194 12195 12196 12197 12198 12199 12200	0000000000	HOH HOH HOH HOH HOH HOH HOH HOH	105 106 107 108 109 110 111 112	51. 853 81. 601 16. 339 1. 00 16. 73 W 59. 699 55. 348 63. 144 1. 00 20. 67 W 45. 186 81. 560 8. 416 1. 00 13. 89 W 37. 516 59. 183 48. 946 1. 00 20. 72 W 22. 032 56. 444 27. 934 1. 00 30. 26 W 65. 773 63. 945 59. 504 1. 00 15. 82 W 45. 931 73. 798 1. 832 1. 00 25. 56 W 29. 602 40. 898 24. 033 1. 00 25. 93 W 19. 080 57. 313 26. 663 1. 00 20. 07 W 61. 355 50. 296 11. 653 1. 00 20. 49 W	0 0 0 0 0 0 0 0

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					FIC	G. 4-	250			(Continued)
ATOM	12201	0	НОН	114	41. 491	58. 601	0.047	1.00 42.91	W	0
ATOM	12202	0	HOH	115	64. 362	64.567	16.259	1.00 24.97	W	0
ATOM	12203	0	HOH	116	43. 928	76.242	2.332	1.00 21.69	W	0
ATOM	12204	0	HOH	117	80. 703	69.349	43.827	1.00 28.64	W	0
ATOM	12205	0	HOH	118	81.671	48.368	20.456	1.00 15.16	W	0
ATOM	12206	0	HOH	119	59. 413	71.127	54.004	1.00 22.01	W	0
ATOM	12207	0	HOH	120	27. 474	69.426	47. 288	1.00 26.74	W	0
ATOM	12208	0	HOH	121	69. 871	60.279	33.380	1.00 13.47	W	0
ATOM	12209	0	HOH	122	67.879	38. 425	47.297	1.00 25.68	W	0
ATOM	12210	0	НОН	123	41.866	62.152	36.306	1.00 27.91	W	. 0
ATOM	12211	0	HOH	124	82.055	50.923	20.718	1.00 23.09	W	0
ATOM	12212	0	HOH	125	38. 821	82.651	33.998	1.00 14.04	W	0
ATOM	12213	0	HOH	126	64. 420	42.195	31.710	1.00 28.88	W	0
ATOM	12214	0	HOH	127	60.713	36. 262	43.885	1.00 22.95	W	0
ATOM	12215	Ō	НОН	128	63.095	38.041	44.744	1.00 26.42	Ÿ	0
ATOM	12216	Ō	НОН	129	36. 718	65. 633	50.633	1.00 38.12	Ÿ	Ö
ATOM	12217	Ö	HOH	130	55. 575	80.086	20. 196	1.00 26.23	W	<u> </u>
ATOM	12218	0	НОН	131	41.981	65. 129	15. 577	1.00 23.62	Ÿ	0
ATOM	12219	Ō	НОН	132	48.067	75.632	53. 563	1.00 36.38	Ÿ	0
ATOM	12220	Ŏ	НОН	133	75.617	59. 792	32.116	1. 00 35. 58	. Ÿ	Ö
ATOM	12221	Ŏ	НОН	134	73. 522	67. 486	30. 484	1.00 21.07	Ÿ	Ö
ATOM	12222	Ŏ	НОН	135	65. 965	81.671	30. 091	1.00 41.74	Ÿ	Ö
ATOM	12223	Ŏ	НОН	136	41.663	53. 300	13.574	1.00 39.95	Ÿ	Ö
ATOM	12224	ŏ	НОН	137	42. 885	39. 029	29.960	1.00 29.57	Ÿ	Ö
ATOM	12225	ŏ	НОН	138	67. 606	56.683	24. 253	1.00 37.19	Ÿ	Ö
ATOM	12226	ŏ	HOH	139	138. 150	54. 591	37. 133	1.00 19.60	W	Ö
ATOM	12227	ŏ	НОН	140	76. 640	48. 505	51.547	1.00 22.87	Ÿ	. 0
ATOM	12228	ŏ	НОН	141	105. 346	35. 319	45. 478	1.00 6.28	Ÿ	Ö
ATOM	12229		НОН	142	108. 946	33. 058	43.850	1.00 17.18	w	. 0
ATOM	12230	ŏ	НОН	143	101. 384	50. 291	32. 321	1.00 12.25	Ÿ	ŏ
ATOM	12231	ŏ	НОН	144	83. 691	56. 732	33. 886	1.00 18.52	Ÿ	Ŏ
ATOM	12232	ŏ	НОН	145	96. 721	59.108	34. 335	1.00 14.59	Ÿ	ŏ
ATOM	12233	ŏ	НОН	146	122. 411	66. 436	57.099	1.00 19.53	Ÿ	ŏ ·
ATOM	12234	ŏ	НОН	147	107. 303	38. 674	48. 678	1.00 12.12	Ÿ	0
ATOM	12235	ŏ	НОН	148	102. 207	54. 174	15. 770	1.00 18.02	Ÿ	ŏ
ATOM	12236	ŏ	НОН	149	104. 534	49. 338	27. 730	1.00 13.02	ẅ	0
ATOM	12237	ŏ	HOH	150	113. 995	67. 497	30. 740	1.00 26.00	Ÿ	ŏ
ATOM	12238	ŏ	НОН	151	115. 903	54. 147	45.005	1.00 20.00	Ÿ	Ö
ATOM	12239	ő	НОН	152	114. 104	55. 650	9. 401	1.00 10.40	Ÿ	0
ATOM	12240	ŏ	НОН	153	86. 360	55. 414	40. 305	1.00 21.03	Ÿ	0
ATOM	12241	ŏ	НОН	154	97. 554	40. 670	45. 200	1.00 14.32	Ÿ	0
ATOM	12242	ő	НОН	155	119. 087	37. 761	27. 531	1.00 10.00	Ψ̈́	
ATOM	12243	0	HOH	156	87. 809	62. 914	36.962	1.00 31.02	W W	0
ATOM	12243	0	HOH	157	83. 356	65. 229	44. 012	1.00 20.29	n W	0
ATOM	12245	0	НОН	158	98. 650					0
ATOM		0	НОН	158		46. 435	54. 377	1.00 26.11	W	0
	12246		нон НОН	160	99. 982	40.104	43.504	1.00 11.71	W	0
ATOM ATOM	12247 12248	0 0	HOH	161	122.550 101.404	42. 243	44.636	1.00 14.84	W	0
ATOM	12249	0	нон НОН	162	88. 481	56. 669 51. 896	35. 498	1.00 35.54 1.00 12.64	W	0
TITOM	14443	U	11011	104	00.401	01.020	31.163	1.00 14.04	H	U

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					E I C	3. 4 <i>-</i>	251			(Continued)
					FIC	J. 4 -	201			
ATOM	12250	0	HOH	163	95.169	58.602	25.005	1.00 10.78	W	0
ATOM	12251	0	НОН	164	115. 235	34.630	45.444	1.00 26.24	W	0 .
ATOM	12252	0	НОН	165	106.826	53.003	55.571	1.00 20.62	W	. 0
ATOM	12253	Õ	НОН	166	84. 875	59. 299	19.482	1.00 36.24	Ÿ	0
ATOM	12254	Ŏ	НОН	167	113. 139	50.670	46. 942	1.00 20.56	Ÿ	Ö
ATOM	12255	Õ	НОН	168	95. 042	48.091	37. 270	1.00 21.34	Ÿ	ŏ
ATOM	12256	ŏ	НОН	169	76. 879	72.537	31.569	1.00 23.37	Ÿ	ő
ATOM	12257	ŏ	НОН	170	114.148	58. 106	48. 086	1.00 18.43	Ÿ	Ö
ATOM	12258	0	НОН	171	89.134	33. 853	32. 584	1.00 22.93	Ÿ	Ö
ATOM	12259	0	НОН	172	104. 484	32. 367	28. 628	1.00 23.01	Ÿ	0
ATOM	12260	0	НОН	173	97. 990	56. 523	56. 950	1.00 25.01	Ÿ	0
ATOM	12261	0	НОН	174	108.093	59.050	11.178	1.00 23.37	W	0
ATOM	12262	0	НОН	175	95.968	47. 759	51. 786	1.00 23.37	W	0
ATOM	12263	0	HOH	176	93. 653	58. 234	55.683	1.00 19.27		
ATOM	12264	0	НОН	177	117.454	64.613	44.832	1.00 19.54	W	0
ATOM	12265	0	НОН						₩	0
ATOM	12266			178	96. 322	67.790	27. 707	1.00 29.36	W	0
		0	НОН	179	80. 831	40.760	23.388	1.00 28.01	W	0
ATOM	12267	0	НОН	180	109. 521	38.188	50. 278	1.00 16.30	W	0
ATOM	12268	0	НОН	181	88. 081	40. 289	29.465	1.00 7.47	W	0
ATOM	12269	0	HOH	182	112. 135	42.102	29.409	1.00 28.14	W	0
ATOM	12270	0	HOH	183	110.546	33. 279	45.877	1.00 22.55	W	0
ATOM	12271	0	НОН	184	101.361	45.858	44.078	1.00 28.83	W	0
ATOM	12272	0	HOH	185	126.633	38. 023	29.778	1.00 31.97	W	0
ATOM	12273	0	HOH	186	122. 283	37. 257	34. 566	1.00 18.77	W	0
ATOM	12274	0	HOH	187	99.753	38. 623	40.032	1.00 18.28	W	0
ATOM ATOM	12275	0	НОН	188	122. 547	56.954	36.341	1.00 20.05	W	0
	12276	0	НОН	189	68.079	78. 219	33. 025	1.00 38.49	W	0
ATOM	12277	0	НОН	190	134.519	46.667		1.00 34.45	W	0
ATOM	12278	0	НОН	191	110. 945	39. 354	35. 865	1.00 10.27	W	0
ATOM	12279	0	НОН	192	118. 982	51.843	57. 881	1.00 13.62	W	0
ATOM	12280	0	НОН	193	123. 824	35.631	32.830	1.00 19.19	W	0
ATOM	12281	0	HOH	194	100. 524	45. 123	38. 393	1.00 26.68	W	0
ATOM	12282	0	НОН	195	122.815	60.696	63. 937	1.00 24.15	W	0
ATOM	12283	0	НОН	196	96. 208	59.856	31.652	1.00 12.71	W	0
ATOM	12284	0	HOH	197	80.023	56. 246	54. 587	1.00 10.61	M	0
ATOM	12285	0	HOH	198	109. 915	41.219	37. 675	1.00 19.28	W	0
ATOM	12286	0	HOH	199	96. 990	75.649	27. 926	1.00 9.03	W	0
ATOM	12287	0	HOH	200	103.494	44. 373	34.046	1.00 8.20	W	0
ATOM	12288	0	НОН	201	97.045	44. 873	53. 124	1.00 15.97	W	0
ATOM	12289	0	НОН	202	109. 135	58. 341	13.499	1.00 22.83	W	0
ATOM	12290	0	HOH	203	96.465	39.089	47.689	1.00 12.68	W	0
ATOM	12291	0	HOH	204	99.669	54. 200	16.885	1.00 13.83	W	0 •
ATOM	12292	0	HOH	205	85.350	34. 351	33. 261	1.00 15.83	W	0
ATOM	12293	0	НОН	206	106. 252	38. 178	46.273	1.00 17.78	W	0
ATOM	12294	0	HOH	207	102. 838	63. 592	15.944	1.00 23.96	W	0
ATOM	12295	0	НОН	208	114. 173	52.027	44.587	1.00 12.16	W	0
ATOM	12296	0	HOH	209	114. 209	49.450	36.803	1.00 19.70	W	0 .
ATOM	12297	0	HOH	210	78. 079	55.141	59.990	1.00 33.63	M	0
ATOM	12298	0	HOH	211	95.004	41.032	14.678	1.00 29.66	W	0

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					FIG	. . 4 -	252			(Continued)
ATOM	12299	0	НОН	212	113.170	36. 816	43. 347	1.00 21.90	W	0
ATOM	12300	0	HOH	213	77. 770	71. 277	45. 572	1.00 31.73	Ÿ	0
ATOM	12301	Õ	НОН	214	128.636	66. 746	61. 783	1.00 37.87	Ÿ	ŏ
ATOM	12302	Õ	НОН	215	128.566	42. 261	18. 644	1.00 26.65	Ÿ	ŏ
ATOM	12303	Ō	НОН	216	135.349	43. 830	34. 280	1.00 24.69	Ÿ	ŏ
ATOM	12304	0	НОН	217	85.640	67.686	27. 706	1.00 32.33	Ÿ	ŏ
ATOM	12305	0	HOH	218	93.669	46.427	45. 506	1.00 24.39	Ÿ	Ö
ATOM	12306	0	HOH	219	117.990	67.819	59.317	1.00 20.28	W	0
ATOM	12307	0	HOH	220	79.954	55.009	62.309	1.00 19.13	W	0
ATOM	12308	0	HOH	221	117. 228	62.083	29.483	1.00 29.50	W	0
ATOM	12309	0	HOH	222	105.505	51.938	31.912	1.00 35.19	W	0
ATOM	12310	0	HOH	223	106.835	57. 215	14.677	1.00 21.77	W	0
ATOM	12311	0	НОН	224	107.489	60.380	64. 395	1.00 24.53	W	0
ATOM	12312	0	НОН	225	79. 753	74. 355	37. 799	1.00 35.35	W	0
ATOM	12313	0	НОН	226	116.807	64. 679	29.466	1.00 24.83	W	0
ATOM	12314	0	НОН	227	87. 239	52. 355	64. 706	1.00 21.19	W	0
ATOM	12315	0	НОН	228	81.916	67. 988	41.878	1.00 14.54	W	0
ATOM	12316	0	НОН	229	106. 295	62. 226	36. 826	1.00 26.06	₩.	0
ATOM ATOM	12317 12318	0	HOH HOH	230 231	78.057	49.553	53. 991	1.00 15.40	W	0
ATOM	12319	0	НОН	232	99. 797 80. 925	47. 673 62. 495	22. 572 37. 326	1.00 18.00	W	0
ATOM	12320	0	НОН	233	93. 378	45. 857	52. 934	1.00 9.28 1.00 12.13	W	0
ATOM	12321	Ö	НОН	234	132.069	46.877	33. 339	1.00 12.13	W	0
ATOM	12322	0	НОН	235	93. 916	62. 211	25. 521	1.00 20.97	W	0 0
ATOM	12323	ŏ	НОН	236	93. 249	60. 882	37. 895	1.00 26.19	W	0
ATOM	12324	Ŏ	НОН	237	100.380	52. 169	18. 636	1.00 20.13	Ϋ́	0
ATOM	12325	Ŏ	НОН	238	82.096	55. 169	32.059	1.00 10.45	Ϋ́	0
ATOM	12326	Õ	НОН	239	94. 471	48. 635	53.699	1.00 13.21	Ÿ	ő
ATOM	12327	0	HOH	240	87.009	55. 227	64. 894	1.00 24.88	Ÿ	ŏ
ATOM	12328	0	HOH	241	95.857	52.760	15.499	1.00 29.83	Ÿ	Ö
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246/10/246

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Ser	Phe	Asp	Gly	Arg	Gly	Ser	Gly	Tyr	Gln	Gly	Asp	Lys	Ile	Met	His
			580					585					590		
Ala	Ile	Asn	Arg	Arg	Leu	Gly	Thr	Phe	Glu	Val	Glu	Asp	Gln	Ile	Glu
		595					600					605			
Ala	Ala	Arg	Gln	Phe	Ser	Lys	Met	Gly	Phe	Val	Asp	Asn	Lys	Arg	Ile
	610					615					620				
Ala	Ile	Trp	Gly	Trp	Ser	Tyr	Gly	Gly	Tyr	Val	Thr	Ser	Met	Val	Leu
625					630					635					640
Gly	Ser	Gly	Ser	Gly	Val	Phe	Lys	Cys	Gly	Ile	Äla	Val	Ala	Pro	Val
				645					650					655	
Ser	Arg	Trp	Glu	Tyr	Tyr	Asp	Ser	Val	Tyr	Thr	Glu	Arg	Tyr	Met	Gly
			660					665					670		
Leu	Pro	Thr	Pro	Glu	Asp	Asn	Leu	Asp	His	Tyr	Arg	Asn	Ser	Thr	Val
		675					680					685			
Met	Ser	Arg	Ala	Glu	Asn	Phe	Lys	Gln	Val	Glu	Tyr	Leu	Leu	Ile	His
	690					695					700				
Gly	Thr	Ala	Asp	Asp	Asn	Val	His	Phe	Gln	Gln	Ser	Ala	Gln	Ile	Ser
705					710					715					720
Lys	Ala	Leu	Val	Asp	Val	Gly	Val	Asp	Phe	Gln	Ala	Met	Trp	Tyr	Thr

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725 730 735

Asp Glu Asp His Gly Ile Ala Ser Ser Thr Ala His Gln His Ile Tyr

740 . 745 75

Thr His Met Ser His Phe Ile Lys Gln Cys Phe Ser Leu Pro

755 760 765

Internation Application No PCT/JP 03/09523

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cliation or other special reason (as specified) cannot be considered to involve an inventive step when the
*O' document referring to an oral disclosure, use, exhibition or other means documents scombined with one or more other such documents, such combination being obvious to a person skilled
"P" document published prior to the International filing date but later than the priority date claimed "&" document member of the same patent family
Date of the actual completion of the international search Date of mailing of the international search report
10 November 2000
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C.(Continu Category °	ation) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	Polovost to shim N
Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Υ .	ABBOTT CATHERINE A ET AL: "Binding to human dipeptidyl peptidase IV by adenosine deaminase and antibodies that inhibit ligand binding involves overlapping, discontinuous sites on a predicted beta propeller domain" EUROPEAN JOURNAL OF BIOCHEMISTRY, vol. 266, no. 3, December 1999 (1999-12), pages 798-810, XP002261851 ISSN: 0014-2956 the whole document	3-5, 14-20
Υ	LAMBEIR A-M ET AL: "A prediction of DPP IV/CD26 domain structure from a physico-chemical investigation of dipeptidyl peptidase IV (CD26) from human seminal plasma" BIOCHIMICA ET BIOPHYSICA ACTA. PROTEIN STRUCTURE AND MOLECULAR ENZYMOLOGY, ELSEVIER, AMSTERDAM,, NL, vol. 1340, no. 2, 18 July 1997 (1997-07-18), pages 215-226, XP004281676 ISSN: 0167-4838 the whole document	3-5, 14-20
Y	MEDRANO F J ET AL: "Structure of proline iminopeptidase from Xanthomonas campestris pv. citri: A prototype for the prolyl oligopeptidase family" EMBO (EUROPEAN MOLECULAR BIOLOGY ORGANIZATION) JOURNAL, vol. 17, no. 1, 2 January 1998 (1998-01-02), pages 1-9, XP002261745 ISSN: 0261-4189 the whole document	3-5, 14-20
Α	POLGAR L: "The prolyl oligopeptidase family" CMLS CELLULAR AND MOLECULAR LIFE SCIENCES, BIRKHAUSER VERLAG, BASEL, CH, vol. 59, no. 2, February 2002 (2002-02), pages 349-362, XP002219152 ISSN: 1420-682X the whole document	1-6, 14-20
A	FULOP V ET AL: "Prolyl oligopeptidase: An unusual beta-propeller domain regulates proteolysis" CELL, CELL PRESS, CAMBRIDGE, NA, US, vol. 94, no. 2, 24 July 1998 (1998-07-24), pages 161-170, XP002221331 ISSN: 0092-8674 the whole document	1-6, 14-20
	the whole document/	

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C.(Continua	tion) DOCUMENTS CONSIDERED TO BE RELEVANT	
Calegory °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
А	AUGUSTYNS K ET AL: "THE UNIQUE PROPERTIES OF DIPEPTIDYL-PEPTIDASE IV (DPP IV/CD26) AND THE THERAPEUTIC POTENTIAL OF DPP IV INHIBITORS" CURRENT MEDICINAL CHEMISTRY, BENTHAM SCIENCE PUBLISHERS BV, BE, vol. 6, no. 4, 1999, pages 311-327, XP000870290 ISSN: 0929-8673 the whole document	1-6, 14-20
P, X	ENGEL MICHAEL ET AL: "The crystal structure of dipeptidyl peptidase IV (CD26) reveals its functional regulation and enzymatic mechanism." PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES, vol. 100, no. 9, 29 April 2003 (2003-04-29), pages 5063-5068, XP002261746 April 29, 2003 ISSN: 0027-8424 (ISSN print) the whole document	1-6, 14-20
Р,Х	RASMUSSEN HANNE B ET AL: "Crystal structure of human dipeptidyl peptidase IV/CD26 in complex with a substrate analog." NATURE STRUCTURAL BIOLOGY, vol. 10, no. 1, January 2003 (2003-01), pages 19-25, XP001168693 ISSN: 1072-8368 (ISSN print) the whole document	1-6, 14-20
Р,Х	HIRAMATSU HAJIME ET AL: "The structure and function of human dipeptidyl peptidase IV, possessing a unique eight-bladed beta-propeller fold." BIOCHEMICAL AND BIOPHYSICAL RESEARCH COMMUNICATIONS, vol. 302, no. 4, 21 March 2003 (2003-03-21), pages 849-854, XP002261748 ISSN: 0006-291X the whole document	1-6, 14-20

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	ition) DOCUMENTS CONSIDERED TO BE RELEVANT	PC1/JP 03/09523
Calegory °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Category	Citation of document, with indication, where appropriate, or the releast personal	Too van to claim No.
P,X	OEFNER CHRISTIAN ET AL: "High-resolution structure of human apo dipeptidyl peptidase IV/CD26 and its complex with 1-'('2-'(5-iodopyridin-2-yl)amino!-ethyl!a mino)- acetyl!-2-cyano-(S)-pyrrolidine." ACTA CRYSTALLOGRAPHICA. SECTION D, BIOLOGICAL CRYSTALLOGRAPHY. DENMARK JUL 2003, vol. 59, no. Pt 7, July 2003 (2003-07), pages 1206-1212, XP008024791 ISSN: 0907-4449 the whole document	1-6, 14-20
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International application No. PCT/JP 03/09523

INTERNATIONAL SEARCH REPORT

Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)			
This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:			
1. X Claims Nos.: 7-13, 22-24 because they relate to subject matter not required to be searched by this Authority, namely:			
see FURTHER INFORMATION sheet PCT/ISA/210			
2. X Claims Nos.: 21 because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:			
see FURTHER INFORMATION sheet PCT/ISA/210			
a Claima Nac			
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).			
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)			
This International Searching Authority found multiple inventions in this international application, as follows:			
1. As all required additional search fees were timely paid by the applicant, this international Search Report covers all searchable claims.			
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.			
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:			
4. No required additional search fees were timely paid by the applicant. Consequently, this international Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:			
Remark on Protest The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.			

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.1

Claims Nos.: 7-13, 22-24

Concerning claims 7 to 13 and 22 to 24 applicant's attention is drawn to Rule 39.1(v) PCT.

The subject-matter of claims 7 to 13 and 22 to 24 refers only to the presentation of structural information and is not regarded as patentable invention within the meaning of Rule 39.1(v) PCT. This information is disclosed e. g. as the atomic coordinates listings (or Tables) of a model, their use in a non-technical method, or said information is stored on a diskette/computer.

Thus, the above mentioned claims will not be searched in accordance with Article 17(2)(a)(i) PCT.

Continuation of Box I.2

Claims Nos.: 21

Present claim 21 relates to a product, i. e. an "effector", defined by reference to a desirable characteristic or property, namely as being an effector of dipeptidyl peptidase IV.

The claim covers all products having this characteristic or property, whereas the application provides no support within the meaning of Article 6 PCT and no disclosure within the meaning of Article 5 PCT of any such products. In the present case, the claim so lacks support, and the application so lacks disclosure, that a meaningful search of the claim is impossible.

Independent of the above reasoning, the claim also lacks clarity (Article 6 PCT). An attempt is made to define the product by reference to a result to be achieved. Again, this lack of clarity in the present case is such as to render a meaningful search over the whole of the claimed scope impossible.

Consequently, no search has been carried out under the provisions of Article 17(2)(a)(ii) PCT.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.